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(54) Title: PHARMACEUTICAL FORMULATIONS COMPRISING AN IMMUNE RESPONSE MODIFIER

(57) Abstract: Pharmaceutical formulations comprising an immune response modifier (IRM) chosen from imidazoquinoline amines, imidazoquinoline amines, imidazopyridine amines, 6,7-fused cycloalkylimidazopyridine amines, 1,2-bridged imidazoquinoline amines, thiazolo-quinolineamines, oxazolo-quinolinamines, thiazolo-pyridinamines, oxazolo-pyridinamines, imidazonaphthyridine amines, tetrahydroimidazonaphthyridine amines, and thiazolonaphthyridine amines; a fatty acid; and a hydrophobic, aprotic component miscible with the fatty acid are useful for the treatment of dermal associated conditions. Novel topical formulations are provided. In one embodiment, the topical formulations are advantageous for treatment of actinic keratosis, postsurgical scars, basal cell carcinoma, atopic dermatitis, and warts.

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PHARMACEUTICAL FORMULATIONS COMPRISING AN IMMUNE RESPONSE MODIFIER

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Field of the Invention

The present invention is directed to pharmaceutical formulations comprising at least one immune response modifier chosen from imidazoquinoline amines, imidazopyridine amines, 6,7-fused cycloalkylimidazopyridine amines, 1,2-bridged imidazoquinoline amines, thiazoloquinoline amines, oxazoloquinoline amines, thiazolopyridine amines, oxazolopyridine amines, imidazonaphthyridine amines, imidazotetrahydronaphthyridine amines, and thiazolonaphthyridine amines. Embodiments of the present invention are directed to topical formulations for application to the skin of a mammal. Other embodiments of the present invention are directed to methods for treating dermal diseases.

Background

Many imidazoquinoline amine, imidazopyridine amine, 6,7-fused cycloalkylimidazopyridine amine, 1,2-bridged imidazoquinoline amine, thiazoloquinoline amine, oxazoloquinoline amine, thiazolopyridine amine, oxazolopyridine amine, imidazonaphthyridine amine, imidazotetrahydronaphthyridine amine, and thiazolonaphthyridine amine compounds have demonstrated potent immunostimulating, antiviral and antitumor (including anticancer) activity, and have also been shown to be useful as vaccine adjuvants. These compounds are hereinafter collectively referred to as "IRM" (immune response modifier) compounds. One of these IRM compounds, known as imiquimod, has been commercialized in a topical formulation, AldaraTM, for the treatment of anogenital warts associated with human papillomavirus.

The mechanism for the antiviral and antitumor activity of these IRM compounds is thought to be due in substantial part to enhancement of the immune response by induction of various important cytokines (e.g., interferons, interleukins, tumor necrosis factor, etc.). Such compounds have been shown to stimulate a rapid release of certain monocyte/macrophage-derived cytokines and are also capable of stimulating B cells to

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secrete antibodies which play an important role in these IRM compounds' antiviral and antitumor activities. One of the predominant immunostimulating responses to these compounds is the induction of interferon (IFN)- α production, which is believed to be very important in the acute antiviral and antitumor activities seen. Moreover, up regulation of other cytokines such as, for example, tumor necrosis factor (TNF), Interleukin-1 (IL-1) and IL-6 also have potentially beneficial activities and are believed to contribute to the antiviral and antitumor properties of these compounds.

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Although some of the beneficial effects of IRMs are known, the ability to provide therapeutic benefit via topical application of an IRM compound for treatment of a particular condition at a particular location may be hindered by a variety of factors. These factors include irritation of the skin to which the formulation is applied, formulation wash away, insolubility and/or degradation of the IRM compound in the formulation, physical instability of the formulation (e.g., separation of components, thickening, precipitation/agglomerization of active ingredient, and the like), poor permeation, and undesired systemic delivery of the topically applied IRM compound. Accordingly, there is a continuing need for new methods and formulations to provide the greatest therapeutic benefit from this class of compounds.

Summary of the Invention

At several locations throughout the specification, guidance is provided through lists of examples. In each instance, the recited list serves only as a representative group; it is not meant that the list is exclusive.

In one aspect, the present invention is directed to a pharmaceutical formulation comprising an immune response modifier selected from imidazoquinoline amines, imidazotetrahydroquinoline amines, imidazopyridine amines, 6,7-fused cycloalkylimidazopyridine amines, 1,2-bridged imidazoquinoline amines, thiazoloquinoline amines, oxazoloquinoline amines, thiazolopyridine amines, oxazolopyridine amines, imidazonaphthyridine amines, imidazotetrahydronaphthyridine amines, and thiazolonaphthyridine amines; a fatty acid; a hydrophobic, aprotic component miscible with the fatty acid and comprising a hydrocarbyl group of 7 or more carbon atoms; and a hydrophilic viscosity enhancing agent selected from cellulose ethers and carbomers.

In one embodiment, the pharmaceutical formulation comprises an immune response modifier selected from imidazonaphthyridine amines, imidazotetrahydronaphthyridine amines, and thiazolonaphthyridine amines; a fatty acid; and a hydrophobic, aprotic component miscible with the fatty acid and comprising a hydrocarbyl group of 7 or more carbon atoms.

The formulation can further comprise one or more of a preservative system, an emulsifier, and water.

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In another aspect, the present invention is directed to a method of treatment of a dermal associated condition comprising applying to skin a topical formulation comprising an immune response modifier selected from imidazoquinoline amines, imidazotetrahydroquinoline amines, imidazopyridine amines, 6,7-fused cycloalkylimidazopyridine amines, 1,2-bridged imidazoquinoline amines, thiazoloquinoline amines, oxazoloquinoline amines, thiazolopyridine amines, oxazolopyridine amines, imidazotetrahydronaphthyridine amines, and thiazolonaphthyridine amines; a fatty acid; a hydrophobic, aprotic component miscible with the fatty acid and comprising a hydrocarbyl group of 7 or more carbon atoms; and a hydrophilic viscosity enhancing agent selected from cellulose ethers and carbomers.

In one embodiment, the method of treatment of a dermal associated condition comprises applying to skin a formulation comprising an immune response modifier selected from imidazonaphthyridine amines, imidazotetrahydronaphthyridine amines, and thiazolonaphthyridine amines; a fatty acid; and a hydrophobic, aprotic component miscible with the fatty acid and comprising a hydrocarbyl group of 7 or more carbon atoms.

In other embodiments, the method of treatment of a dermal associated condition comprises applying to skin a formulation comprising an immune response modifier selected from imidazonaphthyridine amines, imidazotetrahydronaphthyridine amines, and thiazolonaphthyridine amines; a fatty acid; a hydrophobic, aprotic component miscible with the fatty acid and comprising a hydrocarbyl group of 7 or more carbon atoms; and further comprising one or more of a preservative system, an emulsifier, and water.

In one embodiment, the dermal associated condition is selected from actinic keratosis, postsurgical scars, basal cell carcinoma, atopic dermatitis, and warts.

In another aspect, the present invention is directed to a method for delivering an immune response modifier to a dermal surface, the method comprising the steps of selecting a formulation comprising a compound selected from imidazoquinoline amines, imidazotetrahydroquinoline amines, imidazopyridine amines, 6,7-fused cycloalkylimidazopyridine amines, 1,2-bridged imidazoquinoline amines, thiazoloquinoline amines, oxazolo-quinoline amines, thiazolopyridine amines, oxazolopyridine amines, imidazotetrahydronaphthyridine amines, and thiazolonaphthyridine amines; a fatty acid; a hydrophobic, aprotic component miscible with the fatty acid and comprising a hydrocarbyl group of 7 or more carbon atoms; and a hydrophilic viscosity enhancing agent selected from cellulose ethers and carbomers; and applying the selected formulation to the dermal surface for a time sufficient to allow the formulation to deliver the IRM to the dermal surface.

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In one embodiment, the selected formulation comprises an immune response modifier selected from imidazonaphthyridine amines, imidazotetrahydronaphthyridine amines, and thiazolonaphthyridine amines; a fatty acid; and a hydrophobic, aprotic component miscible with the fatty acid and comprising a hydrocarbyl group of 7 or more carbon atoms.

Unless otherwise indicated, all numbers expressing quantities, ratios, and numerical properties of ingredients, reaction conditions, and so forth used in the specification and claims are to be understood as being modified in all instances by the term "about".

As used herein, "a" or "an" or "the" are used interchangeably with "at least one", to mean "one or more" of the element being modified.

Detailed Description

In one aspect, the present invention is directed to a formulation comprising an immune response modifier compound selected from imidazoquinoline amines, imidazotetrahydroquinoline amines, imidazopyridine amines, 6,7-fused cycloalkylimidazopyridine amines, 1,2-bridged imidazoquinoline amines, thiazoloquinoline amines, oxazoloquinoline amines, thiazolopyridine amines, oxazolopyridine amines, imidazotetrahydronaphthyridine amines, and thiazolonaphthyridine amines; a fatty acid; a hydrophobic, aprotic component

miscible with the fatty acid and comprising a hydrocarbyl group of 7 or more carbon atoms, and a hydrophilic viscosity enhancing agent selected from cellulose ethers and carbonners.

These immune response modifier compounds, methods of making them, methods of using them and compositions containing them are disclosed in U.S. Patent Nos. 4,689,338; 4,929,624; 4,988,815; 5,037,986; 5,175,296; 5,238,944; 5,266,575; 5,268,376; 5,346,905; 5,352,784; 5,367,076; 5,389,640; 5,395,937; 5,446,153; 5,482,936; 5,693,811; 5,741,908; 5,756,747; 5,939,090; 6,039,969; 6,083,505; 6,110,929; 6,194,425; 6,245,776; 6,331,539; 6,376,669; and 6,451,810; European Patent 0 394 026; US Publication 2002/0055517; and PCT Publications WO 00/47719; WO 00/76518; WO 01/74343; WO 02/46188; WO 02/46189; WO 02/46190; WO 02/46191; WO 02/46192; WO 02/46193; WO 02/46194; and WO 02/46749 the disclosures of which are incorporated by reference herein.

As noted above, many of the IRM compounds useful in the present invention have demonstrated significant immunomodulating activity. In certain embodiments of the present invention, the IRM compound can be chosen from imidazoquinoline amines, for example, 1*H*-imidazo[4,5-*c*]quinolin-4-amines defined by one of Formulas I-V below:

$$(R_1)_n$$
 N
 R_{21}
 R_{11}

Ι

wherein

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R₁₁ is chosen from alkyl of one to ten carbon atoms, hydroxyalkyl of one to six carbon atoms, acyloxyalkyl wherein the acyloxy moiety is alkanoyloxy of two to four carbon atoms or benzoyloxy, and the alkyl moiety contains one to six carbon atoms, benzyl, (phenyl)ethyl and phenyl, said benzyl, (phenyl)ethyl or phenyl substituent being optionally substituted on the benzene ring by one or two moieties independently chosen from alkyl of one to four carbon atoms, alkoxy of one to four carbon atoms and halogen,

with the proviso that if said benzene ring is substituted by two of said moieties, then said moieties together contain no more than six carbon atoms;

R₂₁ is chosen from hydrogen, alkyl of one to eight carbon atoms, benzyl, (phenyl)ethyl and phenyl, the benzyl, (phenyl)ethyl or phenyl substituent being optionally substituted on the benzene ring by one or two moieties independently chosen from alkyl of one to four carbon atoms, alkoxy of one to four carbon atoms and halogen, with the proviso that when the benzene ring is substituted by two of said moieties, then the moieties together contain no more than six carbon atoms; and

each R_1 is independently chosen from alkoxy of one to four carbon atoms, halogen, and alkyl of one to four carbon atoms, and n is an integer from 0 to 2, with the proviso that if n is 2, then said R_1 groups together contain no more than six carbon atoms;

$$R_{22}$$

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wherein

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R₁₂ is chosen from straight chain or branched chain alkenyl containing two to ten carbon atoms and substituted straight chain or branched chain alkenyl containing two to ten carbon atoms, wherein the substituent is chosen from straight chain or branched chain alkyl containing one to four carbon atoms and cycloalkyl containing three to six carbon atoms; and cycloalkyl containing three to six carbon atoms substituted by straight chain or branched chain alkyl containing one to four carbon atoms; and

R₂₂ is chosen from hydrogen, straight chain or branched chain alkyl containing one to eight carbon atoms, benzyl, (phenyl)ethyl and phenyl, the benzyl, (phenyl)ethyl or phenyl substituent being optionally substituted on the benzene ring by one or two moieties independently chosen from straight chain or branched chain alkyl containing one to four carbon atoms, straight chain or branched chain alkoxy containing one to four carbon

atoms, and halogen, with the proviso that when the benzene ring is substituted by two such moieties, then the moieties together contain no more than six carbon atoms; and

each R₂ is independently chosen from straight chain or branched chain alkoxy containing one to four carbon atoms, halogen, and straight chain or branched chain alkyl containing one to four carbon atoms, and n is an integer from zero to 2, with the proviso that if n is 2, then said R₂ groups together contain no more than six carbon atoms;

$$(R_3)_n$$
 NH_2
 N
 R_{23}

Ш

wherein

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R₂₃ is chosen from hydrogen, straight chain or branched chain alkyl of one to eight carbon atoms, benzyl, (phenyl)ethyl and phenyl, the benzyl, (phenyl)ethyl or phenyl substituent being optionally substituted on the benzene ring by one or two moieties independently chosen from straight chain or branched chain alkyl of one to four carbon atoms, straight chain or branched chain alkoxy of one to four carbon atoms, and halogen, with the proviso that when the benzene ring is substituted by two such moieties, then the moieties together contain no more than six carbon atoms; and

each R_3 is independently chosen from straight chain or branched chain alkoxy of one to four carbon atoms, halogen, and straight chain or branched chain alkyl of one to four carbon atoms, and n is an integer from zero to 2, with the proviso that if n is 2, then said R_3 groups together contain no more than six carbon atoms;

$$R_{4}$$
 IV

wherein

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 R_{14} is -CHR_xR_y wherein R_y is hydrogen or a carbon-carbon bond, with the proviso that when R_y is hydrogen R_x is alkoxy of one to four carbon atoms, hydroxyalkoxy of one to four carbon atoms, 1-alkynyl of two to ten carbon atoms, tetrahydropyranyl, alkoxyalkyl wherein the alkoxy moiety contains one to four carbon atoms and the alkyl moiety contains one to four carbon atoms, or 2-, 3-, or 4-pyridyl, and with the further proviso that when R_y is a carbon-carbon bond R_y and R_x together form a tetrahydrofuranyl group optionally substituted with one or more substituents independently chosen from hydroxy and hydroxyalkyl of one to four carbon atoms;

R₂₄ is chosen from hydrogen, alkyl of one to four carbon atoms, phenyl, and substituted phenyl wherein the substituent is chosen from alkyl of one to four carbon atoms, alkoxy of one to four carbon atoms, and halogen; and

R₄ is chosen from hydrogen, straight chain or branched chain alkoxy containing one to four carbon atoms, halogen, and straight chain or branched chain alkyl containing one to four carbon atoms;

wherein

R₁₅ is chosen from: hydrogen; straight chain or branched chain alkyl containing one to ten carbon atoms and substituted straight chain or branched chain alkyl containing

one to ten carbon atoms, wherein the substituent is chosen from cycloalkyl containing three to six carbon atoms and cycloalkyl containing three to six carbon atoms substituted by straight chain or branched chain alkyl containing one to four carbon atoms; straight chain or branched chain alkenyl containing two to ten carbon atoms and substituted straight chain or branched chain alkenyl containing two to ten carbon atoms, wherein the substituent is chosen from cycloalkyl containing three to six carbon atoms and cycloalkyl containing three to six carbon atoms substituted by straight chain or branched chain alkyl containing one to four carbon atoms; hydroxyalkyl of one to six carbon atoms; alkoxyalkyl wherein the alkoxy moiety contains one to four carbon atoms and the alkyl moiety contains one to six carbon atoms; acyloxyalkyl wherein the acyloxy moiety is alkanoyloxy of two to four carbon atoms or benzoyloxy, and the alkyl moiety contains one to six carbon atoms; benzyl; (phenyl)ethyl; and phenyl; said benzyl, (phenyl)ethyl or phenyl substituent being optionally substituted on the benzene ring by one or two moieties independently chosen from alkyl of one to four carbon atoms, alkoxy of one to four carbon atoms, and halogen, with the proviso that when said benzene ring is substituted by two of said moieties, then the moieties together contain no more than six carbon atoms;

R₂₅ is

$$X$$
 R_s R_T

wherein

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R_S and R_T are independently chosen from hydrogen, alkyl of one to four carbon atoms, phenyl, and substituted phenyl wherein the substituent is chosen from alkyl of one to four carbon atoms, alkoxy of one to four carbon atoms, and halogen;

X is chosen from alkoxy containing one to four carbon atoms, alkoxyalkyl wherein the alkoxy moiety contains one to four carbon atoms and the alkyl moiety contains one to four carbon atoms, hydroxyalkyl of one to four carbon atoms, haloalkyl of one to four carbon atoms, alkylamido wherein the alkyl group contains one to four carbon atoms, amino, substituted amino wherein the substituent is alkyl or hydroxyalkyl of one to four carbon atoms, azido, chloro, hydroxy, 1-morpholino, 1-pyrrolidino, alkylthio of one to four carbon atoms; and

R₅ is chosen from hydrogen, straight chain or branched chain alkoxy containing one to four carbon atoms, halogen, and straight chain or branched chain alkyl containing one to four carbon atoms;

and a pharmaceutically acceptable salt of any of the foregoing.

The IRM compound can also be chosen from 6,7 fused cycloalkylimidazopyridine amines defined by Formula VI below:

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wherein m is 1, 2, or 3;

R₁₆ is chosen from hydrogen; cyclic alkyl of three, four, or five carbon atoms; straight chain or branched chain alkyl containing one to ten carbon atoms and substituted straight chain or branched chain alkyl containing one to ten carbon atoms, wherein the substituent is chosen from cycloalkyl containing three to six carbon atoms and cycloalkyl containing three to six carbon atoms substituted by straight chain or branched chain alkyl containing one to four carbon atoms; fluoro- or chloroalkyl containing from one to ten carbon atoms and one or more fluorine or chlorine atoms; straight chain or branched chain alkenyl containing two to ten carbon atoms and substituted straight chain or branched chain alkenyl containing two to ten carbon atoms, wherein the substituent is chosen from cycloalkyl containing three to six carbon atoms and cycloalkyl containing three to six

carbon atoms substituted by straight chain or branched chain alkyl containing one to four carbon atoms; hydroxyalkyl of one to six carbon atoms; alkoxyalkyl wherein the alkoxy moiety contains one to four carbon atoms and the alkyl moiety contains one to six carbon atoms; acyloxyalkyl wherein the acyloxy moiety is alkanoyloxy of two to four carbon atoms or benzoyloxy, and the alkyl moiety contains one to six carbon atoms, with the proviso that any such alkyl, substituted alkyl, alkenyl, substituted alkenyl, hydroxyalkyl, alkoxyalkyl, or acyloxyalkyl group does not have a fully carbon substituted carbon atom bonded directly to the nitrogen atom; benzyl; (phenyl)ethyl; and phenyl; said benzyl, (phenyl)ethyl or phenyl substituent being optionally substituted on the benzene ring by one or two moieties independently chosen from alkyl of one to four carbon atoms, alkoxy of one to four carbon atoms, and halogen, with the proviso that when said benzene ring is substituted by two of said moieties, then the moieties together contain no more than six carbon atoms;

and -CHR_xR_y

wherein

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 R_y is hydrogen or a carbon-carbon bond, with the proviso that when R_y is hydrogen R_x is alkoxy of one to four carbon atoms, hydroxyalkoxy of one to four carbon atoms, 1-alkynyl of two to ten carbon atoms, tetrahydropyranyl, alkoxyalkyl wherein the alkoxy moiety contains one to four carbon atoms and the alkyl moiety contains one to four carbon atoms, or 2-, 3-, or 4-pyridyl, and with the further proviso that when R_y is a carbon-carbon bond R_y and R_x together form a tetrahydrofuranyl group optionally substituted with one or more substituents independently chosen from hydroxy and hydroxyalkyl of one to four carbon atoms,

R₂₆ is chosen from hydrogen, straight chain or branched chain alkyl containing one to eight carbon atoms, straight chain or branched chain hydroxyalkyl containing one to six carbon atoms, morpholinoalkyl, benzyl, (phenyl)ethyl and phenyl, the benzyl, (phenyl)ethyl or phenyl substituent being optionally substituted on the benzene ring by a moiety chosen from methyl, methoxy, and halogen; and

-C(R_S)(R_T)(X) wherein R_S and R_T are independently chosen from hydrogen, alkyl of one to four carbon atoms, phenyl, and substituted phenyl wherein the substituent is chosen from alkyl of one to four carbon atoms, alkoxy of one to four carbon atoms, and halogen;

X is chosen from alkoxy containing one to four carbon atoms, alkoxyalkyl wherein the alkoxy moiety contains one to four carbon atoms and the alkyl moiety contains one to four carbon atoms, haloalkyl of one to four carbon atoms, alkylamido wherein the alkyl group contains one to four carbon atoms, amino, substituted amino wherein the substituent is alkyl or hydroxyalkyl of one to four carbon atoms, azido, alkylthio of one to four carbon atoms, and morpholinoalkyl wherein the alkyl moiety contains one to four carbon atoms, and

R₆ is chosen from hydrogen, fluoro, chloro, straight chain or branched chain alkyl containing one to four carbon atoms, and straight chain or branched chain fluoro- or chloroalkyl containing one to four carbon atoms and at least one fluorine or chlorine atom; and pharmaceutically acceptable salts thereof.

In other embodiments of the present invention, the IRM compound can be chosen from imidazopyridine amines defined by Formula VII below:

$$\begin{array}{c|c}
NH_{2} \\
N \\
R_{67} \\
R_{77} \\
N \\
R_{17}
\end{array}$$
VII

wherein

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R₁₇ is chosen from hydrogen; -CH₂R_W wherein R_W is chosen from straight chain, branched chain, or cyclic alkyl containing one to ten carbon atoms, straight chain or branched chain alkenyl containing two to ten carbon atoms, straight chain or branched chain hydroxyalkyl containing one to six carbon atoms, alkoxyalkyl wherein the alkoxy moiety contains one to four carbon atoms and the alkyl moiety contains one to six carbon atoms, and phenylethyl; and -CH=CR_ZR_Z wherein each R_Z is independently straight chain, branched chain, or cyclic alkyl of one to six carbon atoms;

R₂₇ is chosen from hydrogen; straight chain or branched chain alkyl containing one to eight carbon atoms; straight chain or branched chain hydroxyalkyl containing one to six carbon atoms; alkoxyalkyl wherein the alkoxy moiety contains one to four carbon atoms and the alkyl moiety contains one to six carbon atoms; benzyl, (phenyl)ethyl and phenyl,

the benzyl, (phenyl)ethyl oand phenyl being optionally substituted on the benzene ring by a moiety chosen from methyl, methoxy, and halogen; and morpholinoalkyl wherein the alkyl moiety contains one to four carbon atoms;

 R_{67} and R_{77} are independently chosen from hydrogen and alkyl of one to five carbon atoms, with the proviso that R_{67} and R_{77} taken together contain no more than six carbon atoms, and with the further proviso that when R_{77} is hydrogen then R_{67} is other than hydrogen and R_{27} is other than hydrogen or morpholinoalkyl, and with the further proviso that when R_{67} is hydrogen then R_{77} and R_{27} are other than hydrogen;

and pharmaceutically acceptable salts thereof.

In yet another embodiment of the present invention, the IRM compound can be chosen from 1,2-bridged imidazoquinoline amines defined by Formula VIII below:

$$(R_8)_q \qquad VIII$$

wherein

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Z is chosen from:

- $(CH_2)_p$ - wherein p is 1 to 4;

-(CH₂)_a-C(R_DR_E)(CH₂)_b-, wherein a and b are integers and a+b is 0 to 3, R_D is hydrogen or alkyl of one to four carbon atoms, and R_E is chosen from alkyl of one to four carbon atoms, hydroxy, -OR_F wherein R_F is alkyl of one to four carbon atoms, and -NR_GR'_G wherein R_G and R'_G are independently hydrogen or alkyl of one to four carbon atoms; and

 $-(CH_2)_a-(Y)-(CH_2)_b$ - wherein a and b are integers and a+b is 0 to 3, and Y is O, S, or -NR_J- wherein R_J is hydrogen or alkyl of one to four carbon atoms;

and wherein q is 0 or 1 and R₈ is chosen from alkyl of one to four carbon atoms, alkoxy of one to four carbon atoms, and halogen,

and pharmaceutically acceptable salts thereof.

In a further embodiment, the IRM compound can be chosen from thiazoloquinoline amines, oxazoloquinoline amines, thiazolonaphthyridine amines, thiazolopyridine amines, and oxazolopyridine amines of Formula IX:

$$R_{39}$$
 R_{49}
 R_{19}
 R_{29}

IX

wherein:

5

R₁₉ is chosen from oxygen, sulfur and selenium;

R₂₉ is chosen from

-hydrogen;

-alkyl;

-alkyl-OH;

-haloalkyl;

-alkenyl;

15 -alkyl-X-alkyl;

-alkyl-X-alkenyl;

-alkenyl-X-alkyl;

-alkenyl-X-alkenyl;

-alkyl-N(R_{59})₂;

20 -alkyl-N₃;

-alkyl-O-C(O)-N(R59)2;

-heterocyclyl;

-alkyl-X-heterocyclyl;

-alkenyl-X-heterocyclyl;

25 -aryl;

-alkyl-X-aryl;

-alkenyl-X-aryl;

-heteroaryl;

-alkyl-X-heteroaryl; and

-alkenyl-X-heteroaryl;

 R_{39} and R_{49} are each independently:

-hydrogen;

-X-alkyl;

-halo;

-haloalkyl;

10 $-N(R_{59})_2$;

or when taken together, R₃₉ and R₄₉ form a fused aromatic, heteroaromatic, cycloalkyl or heterocyclic ring;

X is chosen from -O-, -S-, $-NR_{59}$ -, -C(O)-, -C(O)O-, -OC(O)-, and a bond;

and

each R₅₉ is independently H or C₁₋₈alkyl;

and pharmaceutically acceptable salts thereof.

In another embodiment, the IRM compound can be chosen from imidazonaphthyridine amines and imidazotetrahydronaphthyridine amines of Formulae X and XI below:

20

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wherein

A is =N-CR=CR-CR=; =CR-N=CR-CR=; =CR-CR=N-CR=; or =CR-CR=CR-N=;

 R_{110} is chosen from:

- hydrogen;

```
substituents chosen from:
                                  -aryl;
                                  -heteroaryl;
                                  -heterocyclyl;
   5
                                  -O-C<sub>1-20</sub> alkyl,
                                  -O-(C_{1-20} \text{ alkyl})_{0-1}-\text{aryl};
                                  -O-(C<sub>1-20</sub> alkyl)<sub>0-1</sub>-heteroaryl;
                                 -O-(C<sub>1-20</sub> alkyl)<sub>0-1</sub>-heterocyclyl;
  10
                                 -CO-O-C<sub>1-20</sub> alkyl;
                                 -S(O)_{0-2}-C_{1-20} alkyl;
                                 -S(O)_{0-2}-(C_{1-20} \text{ alkyl})_{0-1}-\text{aryl};
                                 -S(O)_{0-2}-(C_{1-20} \text{ alkyl})_{0-1}-\text{heteroaryl};
                                 -S(O)_{0-2} -(C_{1-20} \text{ alkyl})_{0-1}-heterocyclyl;
 15
                                 -N(R_{310})_2;
                                 -N_3;
                                 oxo;
                                 -halogen;
                                 -NO<sub>2</sub>;
 20
                                 -OH; and
                                -SH; and
                      -C1-20 alkyl-NR310-Q-X-R410 or -C2-20 alkenyl-NR310-Q-X-R410 wherein Q is -CO-
           or -SO_2-; X is a bond, -O- or -NR<sub>310</sub>- and R<sub>410</sub> is aryl; heteroaryl; heterocyclyl; or -C<sub>1-20</sub>
           alkyl or C<sub>2-20</sub> alkenyl that is unsubstituted or substituted by one or more substituents
25
           chosen from:
                                -aryl;
                                -heteroaryl;
                                -heterocyclyl;
                                -O-C<sub>1-20</sub> alkyl,
30
                                -O-(C_{1-20} \text{ alkyl})_{0-1}-aryl;
                                -O-(C_{1-20} \text{ alkyl})_{0-1}-heteroaryl;
```

-C₁₋₂₀ alkyl or C₂₋₂₀ alkenyl that is unsubstituted or substituted by one or more

 $-O-(C_{1-20} \text{ alkyl})_{0-1}$ -heterocyclyl;

-CO-O- C_{1-20} alkyl;

 $-S(O)_{0-2}-C_{1-20}$ alkyl;

 $-S(O)_{0\text{-}2}-\!(C_{1\text{-}20}\;\text{alkyl})_{0\text{-}1}\text{-aryl};$

 $-S(O)_{0-2}-(C_{1-20} \text{ alkyl})_{0-1}$ -heteroaryl;

 $-S(O)_{0-2}$ – $(C_{1-20}$ alkyl $)_{0-1}$ -heterocyclyl;

 $-N(R_{310})_2;$

-NR $_{310}$ -CO-O-C $_{1-20}$ alkyl;

-N₃;

oxo;

10 -halogen;

-NO₂;

-OH; and

-SH; or R_{410} is

15

20

25

wherein Y is -N- or -CR-;

R₂₁₀ is chosen from:

-hydrogen;

-C₁₋₁₀ alkyl;

-C₂₋₁₀ alkenyl;

-aryl;

-C₁₋₁₀ alkyl -O-C₁₋₁₀ alkyl;

-C₁₋₁₀ alkyl-O-C₂₋₁₀ alkenyl; and

-C₁₋₁₀ alkyl or C₂₋₁₀ alkenyl substituted by one or more substituents chosen from:

-OH;

-halogen;

 $-N(R_{310})_2;$

-CO-N(R₃₁₀)₂;

-CO-C₁₋₁₀ alkyl;

 $-N_3$;

-aryl;

-heteroaryl;

-heterocyclyl;

-CO-aryl; and

-CO-heteroaryl;

each R_{310} is independently chosen from hydrogen and C_{1-10} alkyl; and each R is independently chosen from hydrogen, C_{1-10} alkyl, C_{1-10} alkoxy, halogen and trifluoromethyl,

and pharmaceutically acceptable salts thereof;

$$\begin{array}{c|c}
NH_2 \\
N \\
R_{211}
\end{array}$$

$$XI$$

15 wherein

5

10

 $\label{eq:Bis-NR-C(R)2$

R₁₁₁ is chosen from:

- hydrogen;

-C₁₋₂₀ alkyl or C₂₋₂₀ alkenyl that is unsubstituted or substituted by one or more substituents chosen from:

-aryl;

-heteroaryl;

-heterocyclyl;

25 -O-C₁₋₂₀ alkyl;

 $-O-(C_{1-20} alkyl)_{0-1}-aryl;$

-O-(C₁₋₂₀ alkyl)₀₋₁-heteroaryl;

```
-O-(C<sub>1-20</sub> alkyl)<sub>0-1</sub>-heterocyclyl;
                                  -CO-O-C<sub>1-20</sub> alkyl;
                                  -S(O)_{0-2}-C_{1-20} alkyl;
                                  -S(O)_{0-2}-(C_{1-20} \text{ alkyl})_{0-1}-\text{aryl};
                                  -S(O)_{0-2} –(C_{1-20} alkyl)<sub>0-1</sub>-heteroaryl;
                                  -S(O)_{0-2} -(C_{1-20} \text{ alkyl})_{0-1}-heterocyclyl;
                                  -N(R<sub>311</sub>)<sub>2</sub>;
                                  -N<sub>3</sub>;
                                  oxo;
                                  -halogen;
10
                                  -NO<sub>2</sub>;
                                  -OH; and
                                  -SH; and
                       -C1-20 alkyl-NR311-Q-X-R411 or -C2-20 alkenyl-NR311-Q-X-R411 wherein Q is -CO-
            or -SO<sub>2</sub>-; X is a bond, -O- or -NR<sub>311</sub>- and R<sub>411</sub> is aryl; heteroaryl; heterocyclyl; or -C<sub>1-20</sub>
15
            alkyl or C<sub>2-20</sub> alkenyl that is unsubstituted or substituted by one or more substituents
            chosen from:
                                  -aryl;
                                  -heteroaryl;
                                  -heterocyclyl;
20
                                  -O-C<sub>1-20</sub> alkyl,
                                 -O-(C_{1-20} alkyl)_{0-1}-aryl;
                                 -O-(C_{1-20} alkyl)<sub>0-1</sub>-heteroaryl;
                                 -O-(C<sub>1-20</sub> alkyl)<sub>0-1</sub>-heterocyclyl;
                                 -CO-O-C_{1-20} alkyl;
25
                                 -S(O)_{0-2}-C_{1-20} alkyl;
                                 -S(O)_{0-2}-(C_{1-20} \text{ alkyl})_{0-1}-\text{aryl};
                                 -S(O)_{0-2} -(C_{1-20} alkyl)<sub>0-1</sub>-heteroaryl;
                                 -S(O)_{0-2} -(C_{1-20} alkyl)<sub>0-1</sub>-heterocyclyl;
                                 -N(R_{311})_2;
30
                                 -NR_{311}-CO-O-C<sub>1-20</sub> alkyl;
                                 -N_3;
```

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oxo;

-halogen;

-NO₂;

-OH; and

5

-SH; or R₄₁₁ is

wherein Y is -N- or -CR-;

 R_{211} is chosen from:

-hydrogen;

10

15

 $-C_{1-10}$ alkyl;

-C₂₋₁₀ alkenyl;

-aryl

- C_{1-10} alkyl - $O-C_{1-10}$ -alkyl;

-C₁₋₁₀ alkyl-O-C₂₋₁₀ alkenyl; and

-C₁₋₁₀ alkyl or C₂₋₁₀ alkenyl substituted by one or more substituents chosen from:

-OH;

-halogen;

 $-N(R_{311})_2;$

-CO-N(R₃₁₁)₂;

20 -CO- C_{1-10} alkyl;

-N₃;

-aryl;

-heteroaryl;

-heterocyclyl;

25 -CO-aryl; and

-CO-heteroaryl;

each R₃₁₁ is independently chosen from hydrogen and C₁₋₁₀ alkyl; and

each R is independently chosen from hydrogen, C_{1-10} alkyl, C_{1-10} alkoxy, halogen and trifluoromethyl,

and pharmaceutically acceptable salts thereof.

In a further embodiment, the IRM compound can be chosen from imidazoquinoline amines and imidazotetrahydroquinoline amines, for example, 1*H*-imidazo[4,5-*c*]quinolin-4-amines and tetrahydro-1*H*-imidazo[4,5-*c*]quinolin-4-amines defined by Formulas XII, XIII and XIV below:

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10

5

wherein

 R_{112} is -alkyl-NR₃₁₂-CO-R₄₁₂ or -alkenyl-NR₃₁₂-CO- R₄₁₂ wherein R₄₁₂ is aryl, heteroaryl, alkyl or alkenyl, each of which may be unsubstituted or substituted by one or more substituents chosen from:

15 -alkyl; -alkenyl; -alkynyl; -(alkyl)₀₋₁-aryl; -(alkyl)₀₋₁-(substituted aryl); -(alkyl)₀₋₁-heteroaryl; 20 -(alkyl)₀₋₁-(substituted heteroaryl); -O-alkyl; -O-(alkyl)₀₋₁-aryl; -O-(alkyl) $_{0-1}$ -(substituted aryl); -O-(alkyl)₀₋₁-heteroaryl; 25 -O-(alkyl) $_{0-1}$ -(substituted heteroaryl); -CO-aryl; -CO-(substituted aryl);

```
-CO-heteroaryl;
                              -CO-(substituted heteroaryl);
                              -COOH;
                              -CO-O-alkyl;
                              -CO-alkyl;
 5
                              -S(O)_{0-2} -alkyl;
                              -S(O)_{0-2} -(alkyl)<sub>0-1</sub>-aryl;
                              -S(O)<sub>0-2</sub> -(alkyl)<sub>0-1</sub>-(substituted aryl);
                              -S(O)_{0-2} –(alkyl)<sub>0-1</sub>-heteroaryl;
10
                              -S(O)<sub>0-2</sub>-(alkyl)<sub>0-1</sub>-(substituted heteroaryl);
                              -P(O)(OR<sub>312</sub>)<sub>2</sub>;
                              -NR<sub>312</sub>-CO-O-alkyl;
                              -N_3;
                              -halogen;
                              -NO<sub>2</sub>;
15
                              -CN;
                              -haloalkyl;
                              -O-haloalkyl;
                              -CO-haloalkyl;
                              -OH;
20
                              -SH; and in the case of alkyl, alkenyl, or heterocyclyl, oxo;
                              or R_{412} is
                                                                  -(C<sub>1-10</sub>alkyl)-NR<sub>312</sub>-(C<sub>1-10</sub>alkyl)-R<sub>512</sub>
25
                    wherein R<sub>512</sub> is an aryl, (substituted aryl), heteroaryl, (substituted heteroaryl),
          heterocyclyl or (substituted heterocyclyl) group;
                    \mathbf{R}_{212} is chosen from:
                              -hydrogen;
                              -alkyl;
```

-alkenyl;

-aryl;

30

```
-(substituted aryl);
                           -heteroaryl;
                           -(substituted heteroaryl);
                           -heterocyclyl;
  5
                           -(substituted heterocyclyl);
                           -alkyl -O-alkyl;
                           -alkyl-O-alkenyl; and
                           -alkyl or alkenyl substituted by one or more substituents chosen from:
                                   -OH;
 10
                                   -halogen;
                                   -N(R_{312})_2;
                                   -CO-N(R<sub>312</sub>)<sub>2</sub>;
                                   -CO-C<sub>1-10</sub> alkyl;
                                   -CO-O-C_{1-10} alkyl;
15
                                   -N_3;
                                   -aryl;
                                   -(substituted aryl);
                                   -heteroaryl;
                                   -(substituted heteroaryl);
20
                                   -heterocyclyl;
                                   -(substituted heterocyclyl);
                                   -CO-aryl; and
                                   -CO-heteroaryl;
                 each R_{312} is independently chosen from hydrogen; C_{1\text{-}10} alkyl-heteroaryl; C_{1\text{-}10}
         alkyl-(substituted heteroaryl); C_{1-10} alkyl-aryl; C_{1-10} alkyl-(substituted aryl) and C_{1-10}
25
         alkyl;
                 v is 0 to 4;
                 and each R_{12} present is independently chosen from C_{1\text{-}10} alkyl, C_{1\text{-}10} alkoxy,
         halogen and trifluoromethyl;
```

30

XIII

wherein

 R_{113} is -alkyl-NR₃₁₃- SO₂ -X-R₄₁₃ or -alkenyl-NR₃₁₃- SO₂ -X-R₄₁₃;

5 X is a bond or -NR₅₁₃-;

R₄₁₃ is aryl, heteroaryl, heterocyclyl, alkyl or alkenyl, each of which may be unsubstituted or substituted by one or more substituents chosen from:

-alkyl;

-alkenyl;

10 -aryl;

-heteroaryl;

-heterocyclyl;

-substituted cycloalkyl;

-substituted aryl;

-substituted heteroaryl;

-substituted heterocyclyl;

-O-alkyl;

-O-(alkyl)₀₋₁-aryl;

-O-(alkyl)₀₋₁-substituted aryl;

20 -O-(alkyl)₀₋₁-heteroaryl;

-O-(aikyl)₀₋₁-substituted heteroaryl;

-O-(alkyl)₀₋₁-heterocyclyl;

-O-(alkyl)₀₋₁-substituted heterocyclyl;

-COOH;

25 -CO-O-alkyl;

-CO-alkyl;

 $-S(O)_{0-2}$ -alkyl;

 $-S(O)_{0-2}$ -(alkyl)₀₋₁-aryl;

...

```
-S(O)<sub>0-2</sub> -(alkyl)<sub>0-1</sub>-substituted aryl;
                              -S(O)_{0-2} -(alkyl)<sub>0-1</sub>-heteroaryl;
                              -S(O)_{0-2} -(alkyl)<sub>0-1</sub>-substituted heteroaryl;
                              -S(O)_{0-2} -(alkyl)<sub>0-1</sub>-heterocyclyl;
                              -S(O)<sub>0-2</sub> -(alkyl)<sub>0-1</sub>-substituted heterocyclyl;
 5
                              -(alkyl)_{0-1}-NR_{313}R_{313};
                               -(alkyl)<sub>0-1</sub>-NR<sub>313</sub>-CO-O-alkyl;
                              -(alkyl)<sub>0-1</sub>-NR<sub>313</sub>-CO-alkyl;
                               -(alkyl)<sub>0-1</sub>-NR<sub>313</sub>-CO-aryl;
                               -(alkyl)<sub>0-1</sub>-NR<sub>313</sub>-CO-substituted aryl;
10
                               -(alkyl)<sub>0-1</sub>-NR<sub>313</sub>-CO-heteroaryl;
                               -(alkyl)<sub>0-1</sub>-NR<sub>313</sub>-CO-substituted heteroaryl;
                               -N<sub>3</sub>;
                               -halogen;
15
                               -haloalkyl;
                               -haloalkoxy;
                               -CO-haloalkyl;
                               -CO-haloalkoxy;
                               -NO<sub>2</sub>;
                               -CN;
20
                               -OH;
                               -SH; and in the case that R413 is alkyl, alkenyl, or heterocyclyl, oxo;
                     R<sub>213</sub> is chosen from:
                               -hydrogen;
                               -alkyl;
25
                               -alkenyl;
                               -aryl;
                               -substituted aryl;
                               -heteroaryl;
30
                               -substituted heteroaryl;
                               - alkyl-O-alkyl;
                               - alkyl-O- alkenyl; and
```

- alkyl or alkenyl substituted by one or more substituents chosen from: -OH; -halogen; $-N(R_{313})_2;$ -CO-N(R₃₁₃)₂; 5 -CO-C₁₋₁₀ alkyl; -CO-O-C₁₋₁₀ alkyl; $-N_3$; -aryl; -substituted aryl; 10 -heteroaryl; -substituted heteroaryl; -heterocyclyl; -substituted heterocyclyl; -CO-aryl; 15 -CO-(substituted aryl); -CO-heteroaryl; and -CO-(substituted heteroaryl); each R₃₁₃ is independently chosen from hydrogen, C₁₋₁₀ alkyl, and when X is a bond R_{313} and R_{413} can combine to form a 3 to 7 membered heterocyclic or substituted 20 heterocyclic ring;

 R_{513} is chosen from hydrogen, C_{1-10} alkyl, and R_{413} and R_{513} can combine to form a 3 to 7 membered heterocyclic or substituted heterocyclic ring;

v is 0 to 4 and each R_{13} present is independently chosen from C_{1-10} alkyl, C_{1-10} alkoxy, halogen and trifluoromethyl;

25

XIV

wherein

5

10

 \mathbb{R}_{114} is -alkyl-NR₃₁₄-CY-NR₅₁₄-X-R₄₁₄ or -alkenyl-NR₃₁₄-CY- NR₅₁₄-X- R₄₁₄ wherein

Y is = O or = S;

X is a bond, -CO- or $-SO_2$ -;

 \mathbb{R}_{414} is aryl, heteroaryl, heterocyclyl, alkyl or alkenyl, each of which may be unsubstituted or substituted by one or more substituents chosen from:

-alkyl;

-alkenyl;

-aryl;

-heteroaryl;

-heterocyclyl;

-substituted aryl;

-substituted heteroaryl;

-substituted heterocyclyl;

-O-alkyl;

 $-O-(alkyl)_{0-1}-aryl;$

-O-(alkyl)₀₋₁-substituted aryl;

-O-(alkyl)₀₋₁-heteroaryl;

-O-(alkyl)₀₋₁-substituted heteroaryl;

-O-(alkyl)₀₋₁-heterocyclyl;

25 -O-(alkyl)₀₋₁-substituted heterocyclyl;

-COOH;

-CO-O-alkyl;

-CO-alkyl;

```
-S(O)<sub>0-2</sub> -alkyl;
                                 -S(O)_{0-2} –(alkyl)<sub>0-1</sub>-aryl;
                                 -S(O)<sub>0-2</sub> -(alkyl)<sub>0-1</sub>-substituted aryl;
                                 -S(O)_{0-2} –(alkyl)<sub>0-1</sub>-heteroaryl;
                                 -S(O)<sub>0-2</sub> -(alkyl)<sub>0-1</sub>-substituted heteroaryl;
  5
                                 -S(O)<sub>0-2</sub> -(alkyl)<sub>0-1</sub>-heterocyclyl;
                                -S(O)<sub>0-2</sub> -(alkyl)<sub>0-1</sub>-substituted heterocyclyl;
                                 -(alkyl)<sub>0-1</sub>-NR<sub>314</sub>R<sub>314</sub>;
                                -(alkyl)_{0-1}-NR<sub>314</sub>-CO-O-alkyl;
                                -(alkyl)<sub>0-1</sub>-NR<sub>314</sub>-CO-alkyl;
 10
                                -(alkyl)<sub>0-1</sub>-NR<sub>314</sub>-CO-aryl;
                                -(alkyl)<sub>0-1</sub>-NR<sub>314</sub>-CO-substituted aryl;
                                -(alkyl)<sub>0-1</sub>-NR<sub>314</sub>-CO-heteroaryl;
                                -(alkyl)<sub>0-1</sub>-NR<sub>314</sub>-CO-substituted heteroaryl;
15
                                -N_3;
                                -halogen;
                                -haloalkyl;
                                -haloalkoxy;
                                -CO-haloalkoxy;
20
                                -NO<sub>2</sub>;
                                -CN;
                                -OH;
                                -SH; and, in the case that R414 is alkyl, alkenyl or heterocyclyl, oxo;
                     with the proviso that when X is a bond R_{414} can additionally be hydrogen;
25
                     R<sub>214</sub> is chosen from:
                                -hydrogen;
                               -alkyl;
                               -alkenyl;
                                -aryl;
                                -substituted aryl;
30
                               -heteroaryl;
                               -substituted heteroaryl;
```

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- alkyl -O-alkyl;
                          -alkyl-O- alkenyl; and
                          -alkyl or alkenyl substituted by one or more substituents chosen from:
                                  -OH;
                                  -halogen;
5
                                  -N(R314)2;
                                  -CO-N(R<sub>314</sub>)<sub>2</sub>;
                                  -CO-C<sub>1-10</sub> alkyl;
                                  -CO-O-C<sub>1-10</sub> alkyl;
                                   -N_3;
10
                                   -aryl;
                                   -substituted aryl;
                                   -heteroaryl;
                                   -substituted heteroaryl;
                                   -heterocyclyl;
15
                                   -substituted heterocyclyl;
                                   -CO-aryl;
                                   -CO-(substituted aryl);
                                   -CO-heteroaryl; and
                                   -CO-(substituted heteroaryl);
20
                  each R<sub>314</sub> is independently chosen from hydrogen and C<sub>1-10</sub> alkyl;
```

R₅₁₄ is chosen from hydrogen, C₁₋₁₀ alkyl, and R₄₁₄ and R₅₁₄ can combine to form a 3 to 7 membered heterocyclic or substituted heterocyclic ring;

v is 0 to 4 and each R_{14} present is independently chosen from C_{1-10} alkyl, C_{1-10} alkoxy, halogen and trifluoromethyl, and pharmaceutically acceptable salts thereof.

χv

5 wherein:

X is -CHR515-, -CHR515-alkyl-, or -CHR515-alkenyl-;

R₁₁₅ is chosen from:

-R₄₁₅-CR₃₁₅-Z-R₆₁₅-alkyl;

-R₄₁₅-CR₃₁₅-Z-R₆₁₅-alkenyl;

-R₄₁₅-CR₃₁₅-Z-R₆₁₅-aryl;

-R₄₁₅-CR₃₁₅-Z-R₆₁₅-heteroaryl;

- R_{415} - CR_{315} -Z- R_{615} -heterocyclyl;

-R₄₁₅--CR₃₁₅--Z-H;

-R₄₁₅-NR₇₁₅-CR₃₁₅-R₆₁₅-alkyl;

-R₄₁₅-NR₇₁₅ -CR₃₁₅-R₆₁₅-alkenyl;

-R₄₁₅-NR₇₁₅-CR₃₁₅-R₆₁₅-aryl;

-R₄₁₅-NR₇₁₅-CR₃₁₅-R₆₁₅-heteroaryl;

-R₄₁₅-NR₇₁₅-CR₃₁₅-R₆₁₅-heterocyclyl; and

-R₄₁₅-NR₇₁₅ -CR₃₁₅-R₈₁₅;

20

10

15

Z is -NR₅₁₅--, -O--, or -S--;

R₂₁₅ is chosen from:

-hydrogen;

-alkyl;

-alkenyl;

25

-aryl;

-heteroaryl;

-heterocyclyl;

-alkyl-Y-alkyl;

-alkyl-Y-alkenyl;

30

-alkyl-Y-aryl; and

5

25

- alkyl or alkenyl substituted by one or more substituents selected from the group consisting of:

- -OH;
- -haiogen;
- $-N(R_{515})_2;$
 - $-CO-N(R_{515})_2;$
 - -CO-C₁₋₁₀ alkyl;
 - -CO-O-C₁₋₁₀ alkyl;
 - $-N_3$;
- 10 -aryl;
 - -heteroaryl;
 - -heterocyclyl;
 - -CO-aryl; and
 - -CO-heteroaryl;

15
$$R_{315}$$
 is =O or =S;

R415 is alkyl or alkenyl, which may be interrupted by one or more

-O- groups;

each R₅₁₅ is independently H or C₁₋₁₀ alkyl;

R₆₁₅ is a bond, alkyl, or alkenyl, which may be interrupted by one or more

20 –O– groups;

 R_{715} is H, C_{1-10} alkyl, arylalkyl, or R_{415} and R_{715} can join together to form a

5 to 7 membered heterocylcic ring;

 R_{815} is H, C_{1-10} alkyl, or R_{715} and R_{815} can join together to form a 5 to 7 membered heterocyclic ring;

Y is -O- or $-S(O)_{0-2}$;

v is 0 to 4; and

each R_{15} present is independently chosen from C_{1-10} alkyl, C_{1-10} alkoxy, hydroxy, halogen and trifluoromethyl;

wherein:

X is -CHR516-, -CHR516-alkyl-, or -CHR516-alkenyl-;

5

10

R₁₁₆ is chosen from:

 $-R_{416}$ — CR_{316} —Z— R_{616} —alkyl;

-R₄₁₆--CR₃₁₆--Z--R₆₁₆--alkenyl;

 $-R_{416}$ – CR_{316} –Z– R_{616} –aryl;

 $-R_{416}$ – CR_{316} –Z– R_{616} —heteroaryl;

 $-R_{416}$ -CR₃₁₆-Z-R₆₁₆-heterocyclyl;

-R₄₁₆-CR₃₁₆-Z-H;

 $-R_{416}-NR_{716}-CR_{316}-R_{616}-alkyl;$

-R416-NR716-CR316-R616-alkenyl;

 $-R_{416}$ $-NR_{716}$ $-CR_{316}$ $-R_{616}$ -aryl;

 $-R_{416}-NR_{716}-CR_{316}-R_{616}-heteroaryl;$

-R₄₁₆-NR₇₁₆-CR₃₁₆-R₆₁₆-heterocyclyl; and

-R₄₁₆-NR₇₁₆-CR₃₁₆-R₈₁₆;

Z is -NR₅₁₆-, -O-, or -S-;

R₂₁₆ is chosen from:

20

15

-hydrogen;

-alkyl;

-alkenyl;

-aryl;

-heteroaryl;

25

-heterocyclyl;

-alkyl-Y-alkyl;

-alkyl-Y-alkenyl;

-alkyl-Y-aryl; and

from:

-OH; -halogen; 5 $-N(R_{516})_2;$ -CO-N(R₅₁₆)₂; -CO-C₁₋₁₀ alkyl; -CO-O-C₁₋₁₀ alkyl; $-N_3$; 10 -aryl; -heteroaryl; -heterocyclyl; -CO-aryl; and -CO-heteroaryl; 15 R_{316} is =0 or =S; R416 is alkyl or alkenyl, which may be interrupted by one or more -O-groups; each R₅₁₆ is independently H or C₁₋₁₀ alkyl;

 R_{716} is H, $C_{1\text{--}10}$ alkyl, arylalkyl, or R_{416} and R_{716} can join together to form a 5 to 7 membered hetercyclic ring;

R₆₁₆ is a bond, alkyl, or alkenyl, which may be interrupted by one or more

- alkyl or alkenyl substituted by one or more substituents chosen

 R_{816} is H or C_{1-10} alkyl; or R_{716} and R_{816} can join together to form a 5 to 7 membered heterocyclic ring;

Y is -O- or $-S(O)_{0-2}$;

20

25

v is 0 to 4; and

-O- groups;

each R_{16} present is independently chosen from C_{1-10} alkyl, C_{1-10} alkoxy, hydroxy, halogen, and trifluoromethyl;

XVII

wherein:

X is –CHR $_{317}$ -, -CHR $_{317}$ -alkyl-, or –CHR $_{317}$ -alkenyl-;

5

R₁₁₇ is chosen from:

-alkenyl;

-aryl; and

-R417-aryl;

R₂₁₇ is chosen from:

10

-hydrogen;

-alkyl;

-alkenyl;

-aryl;

-heteroaryl;

15

-heterocyclyl;

-alkyl-Y-alkyl;

-alkyl-Y-alkenyl;

-alkyl-Y-aryl; and

- alkyl or alkenyl substituted by one or more substituents chosen

from:

20

25

-OH;

-halogen;

 $-N(R_{317})_2;$

-CO-N(R₃₁₇)₂;

-CO-C₁₋₁₀ alkyl;

-CO-O- C_{1-10} alkyl;

-N₃;

-aryl;

-heteroaryl;

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-heterocyclyl;

-CO-aryl; and

-CO-heteroaryl;

R417 is alkyl or alkenyl, which may be interrupted by one or more

-O- groups;

each R₃₁₇ is independently H or C₁₋₁₀ alkyl;

each Y is independently -O- or -S(O)0-2-;

v is 0 to 4; and

each R_{17} present is independently chosen from C_{1-10} alkyl, C_{1-10} alkoxy,

hydroxy, halogen and trifluoromethyl;

-

15 wherein:

X is -CHR318-, -CHR318-alkyl-, or -CHR318-alkenyl-;

 \mathbf{R}_{118} is chosen from:

-aryl;

-alkenyl; and

-R₄₁₈-aryl;

20

5

10

R₂₁₈ is chosen from:

-hydrogen;

-alkyl;

-alkenyl;

-aryl;

25

-heteroaryl;

-heterocyclyl;

-alkyl-Y-alkyl;

-alkyl-Y-aryl;

- alkyl-Y- alkenyl; and

- alkyl or alkenyl substituted by one or more substituents chosen from:

-OH;

-halogen;

-N(R₃₁₈)₂;

-CO-N(R₃₁₈)₂;

-CO-C₁₋₁₀ alkyl;

-CO-O-C₁₋₁₀ alkyl;

-N₃;

-aryl;

-heteroaryl;

-heterocyclyl;

-CO-aryl; and

-CO-heteroaryl;

R418 is alkyl or alkenyl, which may be interrupted by one or more

-O- groups;

each R₃₁₈ is independently H or C₁₋₁₀ alkyl;

each Y is independently -O- or -S(O)0-2-;

v is 0 to 4; and

each \mathbf{R}_{18} present is independently chosen from $\mathbf{C}_{1\text{-}10}$ alkyl, $\mathbf{C}_{1\text{-}10}$ alkoxy,

hydroxy, halogen and trifluoromethyl;

XIX

25 .

10

15

20

wherein: X is -CHR319-, -CHR319-alkyl-, or -CHR319-alkenyl-;

R₁₁₉ is chosen from:

-heteroaryl;

-heterocyclyl;

```
-R419- heteroaryl; and
                                   -R<sub>419</sub>-heterocyclyl;
                          R_{219} is chosen from:
                                   -hydrogen;
                                   -alkyl;
 5
                                   -alkenyl;
                                   -aryl;
                                   -heteroaryl;
                                   -heterocyclyl;
                                   -alkyl-Y-alkyl;
10
                                   -alkyl-Y-alkenyl;
                                   -alkyl-Y-aryl; and
                                   - alkyl or alkenyl substituted by one or more substituents chosen
                                   from:
                                            -OH;
15
                                            -halogen;
                                            -N(R<sub>319</sub>)<sub>2</sub>;
                                            -CO-N(R<sub>319</sub>)<sub>2</sub>;
                                            -CO-C<sub>1-10</sub> alkyl;
                                            -CO-O-C<sub>1-10</sub> alkyl;
20
                                            -N_3;
                                            -aryl;
                                            -heteroaryl;
                                            -heterocyclyl;
                                            -CO-aryl; and
25
                                            -CO-heteroaryl;
                           R419 is alkyl or alkenyl, which may be interrupted by one or more
                           -O- groups;
                           each R<sub>319</sub> is independently H or C<sub>1-10</sub> alkyl;
                           each Y is independently -O- or -S(O)_{0-2};
30
                           v is 0 to 4; and
```

each R_{19} present is independently chosen from C_{1-10} alkyl, C_{1-10} alkoxy, hydroxy, halogen and trifluoromethyl;

5

wherein:

X is -CHR₃₂₀-, -CHR₃₂₀-alkyl-, or -CHR₃₂₀-alkenyl-;

R₁₂₀ is chosen from:

-heteroaryl;

10

-heterocyclyl;

-R₄₂₀-- heteroaryl; and

-R₄₂₀-heterocyclyl;

 \mathbf{R}_{220} is chosen from:

-hydrogen;

15

-alkyl;

-alkenyl;

-aryl;

-heteroaryl;

-heterocyclyl;

20

-alkyl-Y-alkyl;

-alkyi-Y-alkenyl;

-alkyl-Y-aryl; and

- alkyl or alkenyl substituted by one or more substituents chosen

from:

25

-OH;

-halogen;

 $-N(R_{320})_2;$

-CO-N(R₃₂₀)₂;

-CO-C₁₋₁₀ alkyl;

-CO-O-C₁₋₁₀ alkyl;

-N₃;

-aryl;

-heteroaryl;

-heterocyclyl;

-CO-aryl; and

-CO-heteroaryl;

R₄₂₀ is alkyl or alkenyl, which may be interrupted by one or more

-O- groups;

each R₃₂₀ is independently H or C₁₋₁₀ alkyl;

each Y is independently -O- or -S(O)0-2-;

v is 0 to 4; and

each R_{20} present is independently chosen from $C_{1\text{--}10}$ alkyl, $C_{1\text{--}10}$ alkoxy,

hydroxy, halogen and trifluoromethyl;

15

10

5

XXI

wherein:

X is -CHR521-, -CHR521-alkyl-, or -CHR521-alkenyl-;

20

25

R₁₂₁ is chosen from:

 $-R_{421}$ — NR_{321} — SO_2 — R_{621} —alkyl;

 $-R_{421}$ — NR_{321} — SO_2 — R_{621} —alkenyl;

 $-R_{421}$ $-NR_{321}$ $-SO_2$ $-R_{621}$ -aryl;

 $-R_{421}$ -NR₃₂₁-SO₂-R₆₂₁-heteroaryl;

 $-R_{421}$ -NR₃₂₁-SO₂-R₆₂₁-heterocyclyl;

 $-R_{421}$ — NR_{321} — SO_2 — R_{721} ;

 $-R_{421}-NR_{321}-SO_2-NR_{521}-R_{621}-alkyl;$

-R₄₂₁-NR₃₂₁-SO₂-NR₅₂₁-R₆₂₁-alkenyl;

 $-R_{421}-NR_{321}-SO_2-NR_{521}-R_{621}-aryl;$

```
-R_{421}-NR<sub>321</sub>-SO<sub>2</sub>-NR<sub>521</sub>-R<sub>621</sub>-heteroaryl;
                                         -R<sub>421</sub>-NR<sub>321</sub>-SO<sub>2</sub>-NR<sub>521</sub>-R<sub>621</sub>-heterocyclyl; and
                                         -R<sub>421</sub>-NR<sub>321</sub>-SO<sub>2</sub>-NH<sub>2</sub>;
                               R<sub>221</sub> is chosen from:
 5
                                         -hydrogen;
                                         -alkyl;
                                         -alkenyl;
                                         -aryl;
                                         -heteroaryl;
                                         -heterocyclyl;
10
                                         -alkyl-Y-alkyl;
                                         -alkyl-Y-alkenyl;
                                         -alkyl-Y-aryl; and
                                         - alkyl or alkenyl substituted by one or more substituents chosen
                                         from:
15
                                                   -OH;
                                                   -halogen;
                                                   -N(R_{521})_2;
                                                   -CO-N(R<sub>521</sub>)<sub>2</sub>;
                                                   -CO-C<sub>1-10</sub> alkyl;
20
                                                   -CO-O-C<sub>1-10</sub> alkyl;
                                                   -N_3;
                                                   -aryl;
                                                   -heteroaryl;
                                                   -heterocyclyl;
25
                                                   -CO-aryl; and
                                                   -CO-heteroaryl;
                               Y is -O- or -S(O)<sub>0-2</sub>-;
                               R<sub>321</sub> is H, C<sub>1-10</sub> alkyl, or arylalkyl;
                               each R421 is independently alkyl or alkenyl, which may be interrupted by
30
                               one or more -O- groups, or R<sub>321</sub> and R<sub>421</sub> can join together to form a 5 to 7
                               membered heterocyclic ring;
```

each R_{521} is independently H, $C_{1\text{--}10}$ alkyl, or $C_{2\text{--}10}$ alkenyl;

 \mathbf{R}_{621} is a bond, alkyl, or alkenyl, which may be interrupted by one or more - O- groups;

 R_{721} is C_{1-10} alkyl, or R_{321} and R_{721} can join together to form a 5 to 7 membered heterocyclic ring;

v is 0 to 4; and

each R_{21} present is independently chosen from C_{1-10} alkyl, C_{1-10} alkoxy, hydroxy, halogen and trifluoromethyl;

XXII

10

wherein:

X is -CHR₅₂₂-, -CHR₅₂₂-alkyl-, or -CHR₅₂₂-alkenyl-;

 \mathbf{R}_{122} is chosen from:

15

$$-R_{422}$$
— NR_{322} — SO_2 — R_{622} — $alkyl;$

$$-R_{422}$$
— NR_{322} — SO_2 — R_{622} —alkenyl;

$$-R_{422}\!-\!NR_{322}\!-\!SO_2\!-\!R_{622}\!-\!heteroaryl;$$

20

$$-R_{422}$$
— NR_{322} — SO_2 — R_{722} ,

$$-R_{422}$$
-NR₃₂₂-SO₂-NR₅₂₂-R₆₂₂-alkyl;

-R₄₂₂-NR₃₂₂-SO₂-NR₅₂₂-R₆₂₂-heteroaryl;

25

-R₄₂₂-NR₃₂₂-SO₂-NR₅₂₂-R₆₂₂-heterocyclyl; and

-R₄₂₂-NR₃₂₂-SO₂-NH₂;

R₂₂₂ is chosen from:

-hydrogen;

-alkyl;

```
-alkenyl;
                                  -aryl;
                                  -heteroaryl;
                                  -heterocyclyl;
                                   -alkyl-Y-alkyl;
5
                                   -alkyl-Y-alkenyl;
                                   -alkyl-Y-aryl; and
                                   - alkyl or alkenyl substituted by one or more substituents chosen
                                   from:
                                            -OH;
10
                                           -halogen;
                                            -N(R_{522})_2;
                                            -CO-N(R<sub>522</sub>)<sub>2</sub>;
                                            -CO-C<sub>1-10</sub> alkyl;
                                            -CO-O-C<sub>1-10</sub> alkyl;
15
                                            -N_3;
                                            -aryl;
                                            -heteroaryl;
                                            -heterocyclyl;
                                            -CO-aryl; and
20
                                            -CO-heteroaryl;
                           Y is -O- or -S(O)_{0-2}-;
                           R_{322} is H, C_{1-10} alkyl, or arylalkyl;
                           each R_{422} is independently alkyl or alkenyl, which may be interrupted by
                           one or more -O- groups, or R<sub>322</sub> and R<sub>422</sub> can join together to form a 5 to 7
25
                           membered heterocyclic ring;
                           each R_{522} is independently H, C_{1-10} alkyl, or C_{2-10} alkenyl;
                           \mathbf{R}_{622} is a bond, alkyl, or alkenyl, which may be interrupted by one or more -
                           O- groups;
                           R_{722} is C_{1\text{--}10} alkyl, or R_{322} and R_{722} can join together to form a 5 to 7
30
                           membered heterocyclic ring;
                            v is 0 to 4; and
```

each R_{22} present is independently chosen from C_{1-10} alkyl, C_{1-10} alkoxy, hydroxy, halogen, and trifluoromethyl;

ШХХ

wherein:

X is -CHR₃₂₃-, -CHR₃₂₃-alkyl-, or -CHR₃₂₃-alkenyl-;

Z is -S-, -SO-, or $-SO_2$ -;

 \mathbf{R}_{123} is chosen from:

10

-alkyl;

-aryl;

-heteroaryl;

-heterocyclyl;

-alkenyl;

15

-R₄₂₃-aryl;

-R₄₂₃-heteroaryl;

-R₄₂₃-heterocyclyl;

R₂₂₃ is chosen from:

-hydrogen;

20

-alkyl;

-alkenyl;

-aryl;

-heteroaryl;

-heterocyclyl;

25

-alkyl-Y-alkyl;

- alkyl-Y- alkenyl;

-alkyl-Y-aryl; and

- alkyl or alkenyl substituted by one or more substituents chosen

from:

-OH;

-halogen;

 $-N(R_{323})_2;$

-CO-N(R₃₂₃)₂;

-CO-C₁₋₁₀ alkyl;

-CO-O-C₁₋₁₀ alkyl;

 $-N_3$;

-aryl;

-heteroaryl;

-heterocyclyl;

-CO-aryl; and

-CO-heteroaryl;

each R_{323} is independently H or $C_{1\text{-}10}$ alkyl; each R_{423} is independently alkyl or alkenyl;

each Y is independently -O- or -S(O)₀₋₂-;

v is 0 to 4; and

each R_{23} present is independently chosen from C_{1-10} alkyl, C_{1-10} alkoxy, hydroxy, halogen and trifluoromethyl;

VIXX

wherein:

X is -CHR₃₂₄-, -CHR₃₂₄-alkyl-, or -CHR₃₂₄-alkenyl-;

Z is -S-, -SO-, or $-SO_2-$;

25

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15

R₁₂₄ is chosen from:

-alkyl;

-aryl;

-heteroaryl;

-heterocyclyl;

```
-alkenyl;
                                    -R<sub>424</sub>--aryl;
                                    -R<sub>424</sub>- heteroaryl; and
                                    -R<sub>424</sub>-heterocyclyl;
                           R<sub>224</sub> is chosen from:
                                    -hydrogen;
                                    -alkyl;
                                    -alkenyl;
                                    -aryl;
                                    -heteroaryl;
10
                                    -heterocyclyl;
                                    -alkyl-Y-alkyl;
                                    - alkyl-Y- alkenyl;
                                    -alkyl-Y-aryl; and
15
                                    - alkyl or alkenyl substituted by one or more substituent chosen
                                    from:
                                             -OH;
                                             -halogen;
                                             -N(R_{324})_2;
                                             -CO-N(R<sub>324</sub>)<sub>2</sub>;
20
                                            -CO-C_{1-10} alkyl;
                                             -CO-O-C<sub>1-10</sub> alkyl;
                                             -N<sub>3</sub>;
                                             -aryl;
                                             -heteroaryl;
25
                                             -heterocyclyl;
                                             -CO-aryl; and
                                             -CO-heteroaryl;
                           each R<sub>324</sub> is independently H or C<sub>1-10</sub> alkyl;
                           each R_{424} is independently alkyl or alkenyl;
30
                           each Y is independently -O- or -S(O)0-2-;
                           v is 0 to 4; and
```

each R_{24} present is independently chosen from C_{1-10} alkyl, C_{1-10} alkoxy, hydroxy, halogen and trifluoromethyl;

wherein:

5

10

15

X is -CHR525-, -CHR525-alkyl-, or -CHR525-alkenyl-;

R₁₂₅ is chosen from:

-R₄₂₅-NR₈₂₅-CR₃₂₅-NR₅₂₅-Z-R₆₂₅-alkyl;

 $-R_{425}\!\!-\!\!NR_{825}\!\!-\!\!CR_{325}\!\!-\!\!NR_{525}\!\!-\!\!Z\!\!-\!\!R_{625}\!\!-\!\!alkenyl;$

 $-R_{425}$ $-NR_{825}$ $-CR_{325}$ $-NR_{525}$ -Z $-R_{625}$ -aryl;

 $-R_{425}$ $-NR_{825}$ $-CR_{325}$ $-NR_{525}$ -Z $-R_{625}$ -heteroaryl;

 $-R_{425}$ -NR₈₂₅-CR₃₂₅--NR₅₂₅-Z-R₆₂₅-heterocyclyl;

 $-R_{425}$ -NR₈₂₅—CR₃₂₅—NR₅₂₅R₇₂₅;

 $-R_{425}$ -NR₈₂₅-CR₃₂₅--NR₉₂₅-Z--R₆₂₅-alkyl;

-R₄₂₅-NR₈₂₅-CR₃₂₅-NR₉₂₅-Z-R₆₂₅-alkenyl;

 $-R_{425}\!\!-\!\!NR_{825}\!\!-\!\!CR_{325}\!\!-\!\!NR_{925}\!\!-\!\!Z\!\!-\!\!R_{625}\!\!-\!\!aryl;$

-R₄₂₅-NR₈₂₅-CR₃₂₅-NR₉₂₅-Z-R₆₂₅-heteroaryl; and

 $-R_{425}\!\!-\!\!NR_{825}\!\!-\!\!CR_{325}\!\!-\!\!NR_{925}\!\!-\!\!Z\!\!-\!\!R_{625}\!\!-\!\!heterocyclyl;$

20

R₂₂₅ is chosen from:

-hydrogen;

-alkyl;

-alkenyl;

-aryl;

25

-heteroaryl;

-heterocyclyl;

-alkyl-Y-alkyl;

-alkyl-Y-alkenyl;

-alkyl-Y-aryl; and

- alkyl or alkenyl substituted by one or more substituents chosen from:

-OH;

-halogen;

 $-N(R_{525})_2;$

-CO-N(R₅₂₅)₂;

-CO-C₁₋₁₀ alkyl;

-CO-O-C₁₋₁₀ alkyl;

 $-N_3$;

-aryl;

10

25

30

-heteroaryl;

-heterocyclyl;

-CO-aryl; and

-CO-heteroaryl;

each R_{325} is =O or =S;

each R_{425} is independently alkyl or alkenyl, which may be interrupted by one or more -O- groups;

each R₅₂₅ is independently H or C₁₋₁₀ alkyl;

 R_{625} is a bond, alkyl, or alkenyl, which may be interrupted by one or more

20 –O– groups;

 R_{725} is H, C_{1-10} alkyl which may be interrupted by a hetero atom, or R_{725} can join with R_{525} to form a 5 to 7 membered heterocyclic ring;

 R_{825} is H, C_{1-10} alkyl, arylalkyl, or R_{425} and R_{825} can join together to form a 5 to 7 membered heterocyclic ring;

 R_{925} is C_{1-10} alkyl which can join together with R_{825} to form a 5 to 7 membered heterocyclic ring;

each Y is independently -O- or -S(O)₀₋₂-;

Z is a bond, -CO-, or $-SO_2-$;

v is 0 to 4; and

each R_{25} present is independently chosen from C_{1-10} alkyl, C_{1-10} alkoxy, hydroxy, halogen and trifluoromethyl;

$$(R_{26})_{v}$$
 NH_{2}
 N
 R_{228}
 $XXVI$

wherein:

X is -CHR526-, -CHR526-alkyl-, or -CHR526-alkenyl-;

5

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15

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25

R₁₂₆ is chosen from:

-R₄₂₆-NR₈₂₆-CR₃₂₆--NR₅₂₆-Z-R₆₂₆-alkyl;

-R₄₂₆-NR₈₂₆-CR₃₂₆--NR₅₂₆-Z-R₆₂₆-alkenyl;

-R₄₂₆-NR₈₂₆-CR₃₂₆-NR₅₂₆-Z-R₆₂₆-aryl;

-R₄₂₆-NR₈₂₆-CR₃₂₆-NR₅₂₆-Z-R₆₂₆-heteroaryl;

 $-R_{426}\!\!-\!\!NR_{826}\!\!-\!\!CR_{326}\!\!-\!\!NR_{526}\!\!-\!\!Z\!\!-\!\!R_{626}\!\!-\!\!heterocyclyl;$

-R₄₂₆---NR₈₂₆---CR₃₂₆---NR₅₂₆R₇₂₆;

 $-R_{426}$ $-NR_{826}$ $-CR_{326}$ $-NR_{926}$ -Z $-R_{626}$ -alkyl;

 $-R_{426}\!\!-\!\!NR_{826}\!\!-\!\!CR_{326}\!\!-\!\!NR_{926}\!\!-\!\!Z\!-\!\!-\!\!R_{626}\!\!-\!\!alkenyl;$

 $-R_{426}$ $-NR_{826}$ $-CR_{326}$ $-NR_{926}$ -Z $-R_{626}$ -aryl;

-R $_{426}$ -NR $_{826}$ -CR $_{326}$ -NR $_{926}$ -Z-R $_{626}$ -heteroaryl; and

 $-R_{426}$ -NR₈₂₆-CR₃₂₆-NR₉₂₆-Z-R₆₂₆-heterocyclyl;

R₂₂₆ is chosen from:

-hydrogen;

-alkyl;

-alkenyl;

-aryl;

-heteroaryl;

-heterocyclyl;

-alkyl-Y-alkyl;

-alkyl-Y-alkenyl;

-alkyl-Y-aryl; and

- alkyl or alkenyl substituted by one or more substituents chosen

from:

-OH;

```
-halogen;
                                            -N(R_{526})_2;
                                            -CO-N(R<sub>526</sub>)<sub>2</sub>;
                                            -CO-C<sub>1-10</sub> alkyl;
                                            -CO-O-C<sub>1-10</sub> alkyl;
5
                                            -N_3;
                                            -aryl;
                                            -heteroaryl;
                                            -heterocyclyl;
                                            -CO-aryl; and
10
                                            -CO-heteroaryl;
                           each R_{326} is =O or =S;
                           each R426 is independently alkyl or alkenyl, which may be interrupted by
                           one or more -O- groups;
                           each R<sub>526</sub> is independently H or C<sub>1-10</sub> alkyl;
15
                           R<sub>626</sub> is a bond, alkyl, or alkenyl, which may be interrupted by one or more
                           -O-groups;
                           R_{726} is H, C_{1-10} alkyl which may be interrupted by a hetero atom, or R_{726}
                           can join with R<sub>526</sub> to form a 5 to 7 membered heterocyclic ring;
                           R_{826} is H, C_{1-10} alkyl, arylalkyl, or R_{426} and R_{826} can join together to form a
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                           5 to 7 membered heterocyclic ring;
                           R<sub>926</sub> is C<sub>1-10</sub> alkyl which can join together with R<sub>826</sub> to form a 5 to 7
                           membered heterocyclic ring;
                           each Y is independently -O- or -S(O)<sub>0-2</sub>-;
                           Z is a bond, -CO-, or -SO_2-;
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                           v is 0 to 4; and
                           each R<sub>26</sub> present is independently chosen from C<sub>1-10</sub> alkyl, C<sub>1-10</sub> alkoxy,
                           hydroxy, halogen, and trifluoromethyl;
          and pharmaceutically acceptable salts of any of the foregoing.
                  In another embodiment, the IRM compound can be chosen from 1H-imidazo[4,5-
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c]pyridin-4-amines compounds defined by Formula XXVII

wherein

X is alkylene or alkenylene;

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Z is a bond, -O-, -S-, or $-NR_{527}$ -;

 \mathbf{R}_{127} is aryl, heteroaryl, heterocyclyl, $\mathbf{C}_{1\text{-}20}$ alkyl or

C₂₋₂₀ alkenyl, each of which may be unsubstituted or substituted by one or more substituents independently chosen from:

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-alkyl;

-alkenyl;

-aryl;

-heteroaryl;

-heterocyclyl;

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-substituted cycloalkyl;

-O-alkyl;

-O-(alkyl)₀₋₁-aryl;

-O-(alkyl)₀₋₁-heteroaryl;

-O-(alkyl)₀₋₁-heterocyclyl;

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-COOH;

-CO-O-alkyl;

-CO-alkyl;

 $-S(O)_{0-2}$ -alkyl;

 $-S(O)_{0-2}$ --(alkyl)₀₋₁-aryl;

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 $-S(O)_{0-2}$ –(alkyl)₀₋₁-heteroaryl;

-S(O)₀₋₂ -(alkyl)₀₋₁-heterocyclyl;

-(alkyl)₀₋₁-N(R_{527})₂;

-(alkyl)₀₋₁-NR₅₂₇-CO-O-alkyl;

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-(alkyl)_{0-1}-NR<sub>527</sub>-CO-alkyl;
                                      -(alkyl)<sub>0-1</sub>-NR<sub>527</sub>-CO-aryl;
                                      -(alkyl)<sub>0-1</sub>-NR<sub>527</sub>-CO-heteroaryl;
                                      -N_3;
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                                      -halogen;
                                      -haloalkyl;
                                      -haloalkoxy;
                                      -CO-haloalkyl;
                                      -CO-haloalkoxy;
                                      -NO<sub>2</sub>;
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                                      -CN;
                                      -OH;
                                      -SH; and in the case of alkyl, alkenyl, and heterocyclyl, oxo;
                            R<sub>227</sub> is chosen from:
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                                     -hydrogen;
                                      -alkyl;
                                     -alkenyl;
                                     -alkyl-O-alkyl;
                                     -alkyl-S-alkyl;
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                                     -alkyl-O-aryl;
                                      -alkyl-S-aryl:
                                     -alkyl-O-alkenyl;
                                     -alkyl-S- alkenyl; and
                                     -alkyl or alkenyl substituted by one or more substituents chosen
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                                     from:
                                              -OH;
                                              -halogen;
                                              -N(R_{527})_2;
                                              -CO-N(R<sub>527</sub>)<sub>2</sub>;
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                                              -CS-N(R_{527})_2;
                                              -SO_2-N(R_{527})_2;
                                              -NR_{527}-CO-C_{1-10} alkyl;
```

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-NR<sub>527</sub>-CS-C<sub>1-10</sub> alkyl;
-NR<sub>527</sub>-SO<sub>2</sub>-C<sub>1-10</sub> alkyl;
-CO-C<sub>1-10</sub> alkyl;
-CO-O-C<sub>1-10</sub> alkyl;
-N<sub>3</sub>;
-aryl;
-heteroaryl;
-heterocyclyl;
-CO-aryl; and
-CO-heteroaryl;
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 R_{327} and R_{427} are independently chosen from hydrogen, alkyl, alkenyl, halogen, alkoxy, amino, alkylamino, dialkylamino and alkylthio;

each R₅₂₇ is independently H or C₁₋₁₀alkyl;

and pharmaceutically acceptable salts thereof.

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As used herein, the terms "alkyl", "alkenyl" and the prefix "alk-" are inclusive of both straight chain and branched chain groups and of cyclic groups, i.e. cycloalkyl and cycloalkenyl. Unless otherwise specified, these groups contain from 1 to 20 carbon atoms, with alkenyl groups containing from 2 to 20 carbon atoms. Preferred groups have a total of up to 10 carbon atoms. Cyclic groups can be monocyclic or polycyclic and preferably have from 3 to 10 ring carbon atoms. Exemplary cyclic groups include cyclopropyl, cyclopropylmethyl, cyclopentyl, cyclohexyl and adamantyl.

The term "haloalkyl" is inclusive of groups that are substituted by one or more halogen atoms, including perfluorinated groups. This is also true of groups that include the prefix "halo-". Examples of suitable haloalkyl groups are chloromethyl, trifluoromethyl, and the like.

The term "aryl" as used herein includes carbocyclic aromatic rings or ring systems. Examples of aryl groups include phenyl, naphthyl, biphenyl, fluorenyl and indenyl. The term "heteroaryl" includes aromatic rings or ring systems that contain at least one ring hetero atom (e.g., O, S, N). Suitable heteroaryl groups include furyl, thienyl, pyridyl, quinolinyl, isoquinolinyl, indolyl, isoindolyl, triazolyl, pyrrolyl, tetrazolyl, imidazolyl, pyrazolyl, oxazolyl, thiazolyl, benzofuranyl, benzothiophenyl, carbazolyl, benzoxazolyl,

pyrimidinyl, benzimidazolyl, quinoxalinyl, benzothiazolyl, naphthyridinyl, isoxazolyl, isothiazolyl, purinyl, quinazolinyl, and so on.

"Heterocyclyl" includes non-aromatic rings or ring systems that contain at least one ring hetero atom (e.g., O, S, N) and includes all of the fully saturated and partially unsaturated derivatives of the above mentioned heteroaryl groups. Exemplary heterocyclic groups include pyrrolidinyl, tetrahydrofuranyl, morpholinyl, thiomorpholinyl, piperidinyl, piperazinyl, thiazolidinyl, imidazolidinyl, isothiazolidinyl, and the like.

In some embodiments, the topical formulations of the present invention are prepared using the free base form of the IRM compound.

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The amount of an IRM compound that will be therapeutically effective in a specific situation will depend on such things as the activity of the particular compound, the dosing regimen, the application site, the particular formulation and the condition being treated. As such, it is generally not practical to identify specific administration amounts herein; however, those skilled in the art will be able to determine appropriate therapeutically effective amounts based on the guidance provided herein, information available in the art pertaining to these compounds, and routine testing. The term "a therapeutically effective amount" means an amount of the compound sufficient to induce a therapeutic effect, such as cytokine induction, inhibition of TH2 immune response, antiviral or antitumor activity, reduction or elimination of postsurgical scarring, or reduction or resolution of actinic keratosis or pre-actinic keratosis lesions.

In general, the amount of the IRM compound present in a topical formulation of the invention will be an amount effective to treat a targeted condition, to prevent recurrence of the condition, or to promote immunity against the condition. The amount or concentration of the IRM compound can range from 0.001% to 10% by weight based on the total formulation weight, such as, for example, from 0.03% to 5.0% by weight, or from 0.1 to 1.0% by weight. In certain embodiments, the amount of the IRM compound is at least 0.003% by weight, such as, for example, at least 0.005%, at least 0.01%, at least 0.03%, at least 0.10%, at least 0.30% and at least 1.0%. In other embodiments, the amount of the IRM compound is at most 5.0% by weight, such as, for example, at most 3.0%, and at most 1.0%.

The topical formulations of the invention additionally comprise a fatty acid. As used herein, the term "fatty acid" means a carboxylic acid, either saturated or unsaturated,

comprising 6 to 28 carbon atoms, such as, for example, from 10 to 22 carbon atoms. Non-limiting examples of such fatty acids include isostearic acid, oleic acid, and linear-or-branched chained carboxylic acids of 6 to 18 carbon atoms. The fatty acid may be present in the formulation in an amount sufficient to solubilize the IRM compound. In one embodiment, the amount of the fatty acid can range from 0.05 % to 40 % by weight based on the total weight of the formulation, such as, for example, from 1% to 30%, from 3% to 15% and from 5% to 10%. In certain embodiments, the amount of the fatty acid is at least 3.0% by weight, such as, for example, at least 5.0%, at least 10.0%, and at least 25%. The fatty acid component of the formulation can comprise one or more fatty acids.

The topical formulations of the invention additionally comprise at least one hydrophobic, aprotic component miscible with the fatty acid and comprising a hydrocarbyl group of 7 or more carbon atoms. By "hydrophobic" is meant that the component is essentially insoluble in water, i.e. immiscible with water and unable to form a micelle in water, and does not contain polyoxyethylene or acid salt groups. Preferably the hydrophobic, aprotic component has a hydrophilic lipophilic balance (HLB) of less than 2. The HLB of a component may be determined as described, for example, in Attwood, D., Florence, A. T. Surfactant Systems: Their Chemistry, Pharmacy, and Biology. New York: Chapman & Hall, 471-473, 1983. By "aprotic" is meant that the component cannot donate a proton to the IRM and does not contain groups such as carboxyl, hydroxy, primary and secondary amino, primary and secondary amido, or quaternary ammonium groups. Preferably this component has a pKa of at least 14.2 and does not substantially solubilize or form a complex such as an acid-base pair or complex or a hydrogen bond complex with the IRM compound. By "not substantially" is meant that the ratio of the IRM compound's solubility in the hydrophilic, aprotic component to that in isostearic acid is less than 1:40.

Formulations intended for dermal or topical use desirably have a certain minimum amount of an oil phase to provide qualities such as spreadability, feel on the skin, texture, and so on. However, if all the components of the oil phase solubilize the IRM, then the degree of saturation of the IRM in the formulation will decrease, making it more difficult to deliver the IRM from the formulation to the skin. Addition of the hydrophobic, aprotic component can increase the oil phase volume of the topical formulation to provide desirable qualitites such as spreadability and feel, while at the same time not appreciably altering the degree of saturation or thermodynamic activity of the IRM. For example, the

amount of fatty acid, which solubilizes the IRM, can be reduced to increase the degree of IRM saturation while maintaining a sufficient oil phase volume by virtue of the addition of the hydrophobic, aprotic component, which does not offset the increased IRM saturation. Thus, the topical formulation of the present invention can facilitate both physical property and drug delivery requirements. Degree of saturation and thermodynamic activity of the IRM in these formulations is equal to the IRM concentration in the oil phase divided by the saturation concentration of the IRM in the oil phase. When the topical formulations of the present invention contain saturated IRM the thermodynamic activity or degree of saturation is unity, and when partially saturated the thermodynamic activity or degree of saturation is less than unity.

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The amount of the hydrophobic, aprotic component present in a formulation of the invention can range from 1% to 30% by weight based on the total formulation weight, for example, from 3 % to 15% by weight, and from 5 to 10% by weight. In certain embodiments, the amount of the hydrophobic, aprotic component is at least 3.0% by weight, for example, at least 5.0%, and at least 10.0%. The weight ratio of the hydrophobic, aprotic component to the fatty acid can be 0.025:1 to 600:1, for example, 0.5:1 to 50:1, and 2:1 to 30:1. The combined amount (weight percent of the total topical formulation weight) of the hydrophobic, aprotic component and the fatty acid can be 2% to 50% by weight, for example 2% to 30%, 5% to 30%, 5% to 20%, and 10% to 20%.

Examples of useful hydrophobic, aprotic components include but are not limited to fatty acid esters, for example, isopropyl mysristate, isopropyl palmitate, diisopropyl dimer dilinoleate; triglycerides, for example, caprylic/capric triglyceride; cetyl esters wax; hydrocarbons of 8 or more carbon atoms, for example, light mineral oil, white petrolatum; and waxes, for example, beeswax. In some embodiments, the hydrophobic, aprotic component is chosen from one or more of isopropyl mysristate, isopropyl palmitate, caprylic/capric triglyceride, and diisopropyl dimer dilinoleate.

The formulations of the present invention can also comprise a hydrophilic viscosity enhancing agent. Examples of suitable hydrophilic viscosity enhancing agents include cellulose ethers such as hydroxypropylmethylcellulose, hydroxyethylcellulose, hydroxypropylcellulose, and carboxymethylcellulose; polysaccharide gums such as xanthan gum; and homopolymers and copolymers of acrylic acid crosslinked with allyl sucrose or allyl pentaerythriol such as those polymers designated as carbomers in the

United States Pharmacopoeia. Suitable carborners include, for example, those available as CarbopolTM 934P, Carbopol 971P, Carbopol 940, Carbopol 974P, Carbopol 980, and PemulenTM TR-1 (USP/NF Monograph; Carbomer 1342), all available from Noveon, Cleveland, Ohio. In one embodiment of the present invention, the viscosity enhancing agent is chosen from Carbopol 974P and 980. When included, the viscosity enhancing agent is generally present in an amount ranging from 0.1% to 10% by weight of total formulation weight, such as, for example, from 0.5 % to 5% by weight, from 0.5% to 1.5% by weight, and from 0.7% to 3% by weight. In certain embodiments, the amount of the viscosity enhancing agent is at least 0.5% by weight, for example, at least 0.6% by weight, at least 0.7% by weight, at least 0.9% by weight, and at least 1.0% by weight.

The formulations of the invention can additionally comprise an emulsifier. Suitable emulsifiers include non-ionic surfactants such as, for example, polysorbate 60, sorbitan monostearate, polyglyceryl-4 oleate, polyoxyethylene(4) lauryl ether, etc. In certain embodiments, the emulsifier is chosen from poloxamers (e.g., Pluronic™ F68, also known as Poloxamer 188, a poly(ethylene glycol)-block-poly(propylene glycol)-block-poly(ethylene glycol), available from BASF, Ludwigshafen, Germany) and sorbitan trioleate (e.g., Span 85 available from Uniqema, New Castle, DE). If included, the emulsifier is generally present in an amount of 0.1% to 10% by weight of total formulation weight, for example, from 0.5% to 5% by weight, and from 0.75% to 3.5% by weight. In certain embodiments, the amount of the emulsifier is at least 1.0% by weight, for example, at least 2.5%, at least 3.5%, and at least 5.0%.

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In certain embodiments of the present invention, the formulation can also include at least one chelating agent. The chelating agent functions to chelate metal ions that may be present in the formulation. Suitable chelating agents include salts of ethylenediaminetetraacetate (EDTA), such as the disodium salt. If included, the chelating agent is generally present in an amount ranging from 0.001 % to 0.1% by weight, and preferably from 0.01% to 0.05% by weight. In certain embodiments, the amount of the chelating agent is at least 0.005% by weight, such as, for example, at least 0.01%, and at least 0.05%.

The formulation can also include a preservative system. The preservative system is generally comprised of at least one preservative compound chosen from methylparaben, ethylparaben, propylparaben, phenoxyethanol, iodopropynyl butylcarbamate, sorbic acid,

a fatty acid monoester of glycerin such as glycerol monolaurate, and a fatty acid monoester of propylene glycol such as propylene glycol monocaprylate. The preservative system may also include a preservative enhancing solubilizer which enhances the solubility of the preservative in the aqueous phase, examples of which include diethylene glycol monoethyl ether and propylene glycol. In one embodiment, the preservative system can be comprised of methylparaben, propylparaben, and propylene glycol. In another embodiment, the preservative system can be comprised of methylparaben, ethylparaben, and diethylene glycol monoethyl ether. In one embodiment, the preservative system can be comprised of phenoxyethanol, methylparaben or methyl- and ethylparaben, and diethylene glycol monoethyl ether. In another embodiment, the preservative system can be comprised of iodopropynyl butylcarbamate. In another embodiment, the preservative system can be comprised of iodopropynyl butylcarbamate, diethylene glycol monoethyl ether, and poly(ethylene glycol)(4) monolaurate. In another embodiment, the preservative system can be comprised of iodopropynyl butylcarbamate, one or more of methylparaben, ethylparaben, propylparaben, or phenoxyethanol, and diethylene glycol monoethyl ether. In the above embodiments, the methylparaben, ethylparaben, and propylparaben can each be present in the formulations in an amount ranging from 0.01% to 0.5% by weight of the formulation weight, for example, from 0.05 % to 0.25% by weight, and from 0.1% to 0.2% by weight. The iodopropynyl butylcarbamate can be present in the formulations in an amount ranging from 0.01% to 0.1%. The phenoxyethanol can be present in the formulations in an amount ranging from 0.1% to 1%. The propylene glycol and diethylene glycol monoethyl ether can each be present in the formulations in an amount ranging from 1% to 30% by weight of the formulation weight, such as, for example, from 5 % to 25% by weight, and from 10% to 15% by weight. The preservative system can be present in the formulations in an amount ranging from 0.01% to 30% by weight of the formulation weight, for example, from 0.05% to 30%, from 0.1% to 25% by weight, and from 0.2% to 15% by weight. In a further embodiment, the methylparaben, ethylparaben, propylparaben, iodopropynyl butylcarbamate, and phenoxyethanol can be solubilized in propylene glycol, poly(ethylene glycol)(4) monolaurate, or diethylene glycol monoethyl ether prior to addition to the formulation. The preservative system can be selected such that it meets the criteria for antimicrobial effectiveness set forth in the United States Pharmacopeia <51>.

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The formulations of the present invention may additionally comprise at least one pH adjuster. Suitable pH adjusters include organic bases and inorganic bases such as, for example, KOH, NaOH. The pH of the topical formulations of the present invention generally ranges from 3.5 to 7.0. In one embodiment, the pH of the topical formulations of the present invention can range from 4.0 to 6.0, preferably 5.0. In another embodiment of the invention, the pH of the topical formulations of the present invention can range from 5.5 to 6.5, preferably 6.0.

Any of the foregoing formulations can be in the form of an oil-in-water emulsion such as a cream or a lotion. Such an emulsion can comprise an oil phase comprising the IRM compounds, a fatty acid in an amount sufficient to solubilize the IRM compounds, a hydrophobic, aprotic component; and an aqueous phase comprising a hydrophilic viscosity enhancing agent, for example, a carbomer. In certain embodiments, the amount or concentration of the IRM in the oil phase can be at least 0.01%, for example, at least 0.02%, at least 0.1%, and at least 1% with respect to oil phase weight. In other embodiments, the amount or concentration of the IRM in the oil phase can be at most 20%, for example, at most 10%, and at most 5% with respect to oil phase weight. The emulsion can be preserved so that when challenged by an antimicrobial effectiveness test, it meets regulatory requirements for topical creams packaged in multiple-use containers.

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Any of the foregoing formulations according to the present invention can be applied to the dermal surfaces of a mammal. Depending on the IRM compound concentration, formulation composition, and dermal surface, the therapeutic effect of the IRM compound may extend only to the superficial layers of the dermal surface or to tissues below the dermal surface. Thus, another aspect of the present invention is directed to a method for the treatment of a dermal associated condition comprising applying to skin one of the foregoing formulations. As used herein, a "dermal associated condition" means an inflammatory, infectious, neoplastic or other condition that involves a dermal surface or that is in sufficient proximity to a dermal surface to be affected by a therapeutic agent topically applied to the dermal surface. Examples of a dermal associated condition include warts, atopic dermatitis, basal cell carcinoma, postsurgical scars, and actinic keratosis.

In one embodiment, the formulations can be applied to the surface of skin for treatment of actinic keratosis (AK). Actinic keratoses are premalignant lesions considered

biologically to be either carcinoma in-situ or squamous intraepidermal neoplasia. AK is the most frequent epidermal tumor and is induced by ultraviolet (UV) radiation, typically from sunlight. Because of its precancerous nature, AK may be considered the most important manifestation of sun-induced skin damage.

In some embodiments, the above described formulations are particularly advantageous for dermal application for a period of time sufficient to obtain a desired therapeutic effect without undesired systemic absorption of the IRM.

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EXAMPLES

The following Examples are provided to further describe various IRM formulations and methods according to the invention. The examples, however, are not intended to limit the formulations and methods within the spirit and scope of the invention.

Examples 1-7 and Comparative Example C1

Table 1 summarizes topical formulations made in accordance with the present invention in a percentage weight-by-weight basis.

TABLE

Ingredient (Compendial Status)			T (percent	Topical Cream (percentage weight-by-weight)	weight)			
	Comparative Example C1 (Placebo)	Example 1	Example 2	Example 3	Example 4	Example 5	Example 6	Example 7
IRM Compound 1	0.00	0.001	0.003	0.010	0.03	0.10	0:30	1.00
Isostearic Acid	5.00	5.00	5.00	5.00	5.00	5.00	7.00	10.00
Isopropyl Myristate (NF)	10.00	10.00	10.00	10.00	10.00	10.00	8.00	5.00
Carbomer 974P (NF)	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00
Poloxamer 188 (NF)	2.50	2.50	2.50	2.50	2.50	2.50	2.50	2.50
Propylene Glycol (USP)	15.00	15.00	15.00	15.00	15.00	15.00	15.00	15.00
Methylparaben (NF)	0.20	0.20	0.20	0.20	0.20	0.20	0.20	0.20
Propylparaben (NF)	0.10	0.10	01.0	0.10	0.10	0.10	0.10	0.10
Edetate Disodium (USP)	0.05	0.05	0.05	0.05	0.05	0.05	0.05	0.05
Sodium Hydroxide (NF) Solution, 20% w/w	0.50	0.50	0.50	0.50	0.50	0.50	0.50	0.55
Purified Water (USP)	65.65	65.649	65.647	65.64	65.62	65.55	65.35	64.60
Total	. 100.00	100.00	100.00	100.00	100.00	100.00	100.00	100.00

The formulations set forth in Table 1 were prepared in the following manner:

Oil phase preparation: 2-methyl-1-(2-methylpropyl)-1H-imidazo[4,5-c]

[1,5]naphthyridin-4-amine (IRM compound 1) was dissolved in isostearic acid and isopropyl myristate, with heat if necessary. Carbomer 974P was then dispersed in the oil phase.

Water phase preparation: Edetate disodium was dissolved in the water.

Methylparaben and propylparaben were dissolved in propylene glycol and the solution was subsequently added to the water phase. Poloxamer 188 was then added to the water phase and mixed until dissolved.

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<u>Phase combination:</u> The oil phase was added to the water phase at ambient conditions. The emulsion was then homogenized. After homogenization, sodium hydroxide solution (20% w/w) was added and the resulting cream was mixed until smooth and uniform. The pH of the cream was measured and a pH adjustment was made with additional sodium hydroxide solution, if necessary, to meet the in-process target pH of 5.

Formulations containing 2-methyl-1-(2-methylpropyl)-1H-imidazo[4,5-c] [1,5]naphthyridin-4-amine (IRM Compound 1) were tested for their ability to induce increases in cytokine concentrations in rats following topical application. This study was undertaken to evaluate cytokine induction following a single dosing of various strengths and timepoints or a multiple vs. single dosing of IRM Compound 1. The formulations described above were tested by examining tissue and serum concentrations of TNF- α , MCP-1 (monocyte chemoattractant protein-1) and IFN- α cytokines following drug treatment.

Female CD hairless rats (Charles River Laboratories, Wilmington, MA) weighing 200-250 grams were used in all studies. Animals were randomized to treatment groups and dosed five per treatment group.

The rats were acclimated to collars around the neck on two consecutive days prior to actual dosing. The rats were collared before dosing to prevent ingestion of the drug, and were then dosed topically with 50 μ L of active cream or the appropriate placebo on right flank and then housed individually following dosing. At various times following dosing, the rats were anesthetized and blood was collected by cardiac puncture. Blood was allowed to clot at room temperature and serum was separated from the clot via centrifugation and stored at -20 °C until it was analyzed for cytokine concentrations.

Following blood collection, the rats were euthanized and their skins removed. Tissue from both treated site (at) and contralateral site (away) were obtained using an 8 mm punch biopsy, weighed, placed in a sealed 1.8 ml cryovial and flash frozen in liquid nitrogen. The frozen tissue sample was then suspended in 1.0 mL RPMI medium (Celox, Hopkins, MN) containing 10% fetal bovine serum (Sigma, St. Louis, MO), 2 mM L-glutamine, penicillin/streptomycin, and 2-mercaptoethanol (RPMI complete) combined with a protease inhibitor cocktail set III (Calbiochem, San Diego, CA). The tissue was homogenized using a Tissue TearorTM (Biospec Products, Bartlesville, OK) for approximately 1 minute. The tissue suspension was then centrifuged at 2000 rpm for 10 minutes under refrigeration to pellet debris, and the supernatant collected and stored at -20 °C until analyzed for cytokine concentrations.

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ELISAs for rat MCP-1 were purchased from BioSource Intl. (Camarillo, CA) and rat TNF-α were purchased from BD Pharmingen (San Diego, CA) and performed according to manufacturer's specifications. Results for both TNF-α, and MCP-1 were expressed in pg/200 mg tissue or pg/ml serum. The sensitivity of the TNF-α ELISA was 31.2 pg/ml and of the MCP-1 ELISA was 11.7 pg/ml. IFN-α concentrations in both serum and skin tissue were determined using a bioassay that measured inhibition of the viral cytopathic effect of vesicular stomatitis virus on rat LMS-C2 fibroblast cells as previously described (Reiter, M. J., Testerman, T. L., Miller, R. L., Weeks, C. E., and Tomai, M. A. (1994) "Cytokine Induction in Mice by the Immunomodulator Imiquimod." J. Leukocyte Biol. 55, 234-240). IIT Research Institute, Chicago IL, performed these assays. Results for IFN-α concentrations were normalized to a standard reference rat IFN-α, preparation with results being reported in U/mL and are normalized per mg of tissue.

The data shown below in Tables 2-4 are from three separate experiments and analyzed to 1) measure pharmacokinetics by full time course, 2) measure dose response and 3) measure multiple vs. single dosing.

In order to determine the kinetics of local and systemic cytokine production following local administration of IRM Compound 1, the full time course study (Study 1 with results in Table 2) was done by topically dosing rats with the topical cream formulation of Example 7.

Serum and tissue samples were taken at 1, 2, 4, 8, 16, 24 and 48 hours post dose. Multiple cytokines (MCP-1, TNF- α and IFN- α) were analyzed separately.

With the tissue data, for each hour measured, a paired t-test (used to eliminate within subject variability) analyzed the difference between treated tissue and control tissue from the same animal. A p-value less than alpha=0.05 indicated a statistically significant difference between the treated and control tissue at that hour. The data are presented in Table 2.

Table 2. Cytokine Concentrations in Rat Serum and Dermal Tissue Following Application of the Topical Formulation of Example 7 Full Time Course^a

		Су	tokine Concentra	ition ^b
Time (hours)			TNF-α	
Post Dose	Dose	Serum	Treated Site	Control site
0	untreated	0	NA ·	96±5
16	placebo	0	103+8	71 <u>+</u> 6
1	1%	6 <u>+</u> 6	318 <u>+</u> 33°	96 <u>+</u> 13
2	1%	0	1125 <u>+</u> 74°	124 <u>+</u> 18
4	1%	0	1120 <u>+</u> 51 °	129±11
8	1%	24 <u>+16</u>	429±56°	91 <u>+</u> 12
16	1%	6 <u>+</u> 4	231 <u>+</u> 22 °	87 <u>+</u> 27
24	1%	32 <u>+</u> 32	198 <u>+</u> 28°	103±13
48	1%	49 <u>+</u> 49	74 <u>+</u> 10	69 <u>+</u> 15
			MCP-1	•
0	untreated	81 <u>+</u> 30	NA	44 <u>+</u> 2
16	placebo	144 <u>+</u> 9	144 <u>+</u> 41	42 <u>+</u> 3
1	1%	86 <u>+</u> 29	40 <u>+</u> 8	42 <u>+</u> 3
2	1%	123 <u>+</u> 31	234 <u>+</u> 29°	50 <u>+</u> 4
4	1%	101 <u>+</u> 28	723 <u>+</u> 89°	41 <u>+</u> 5
8	1%	438 <u>+</u> 91 °	1474±202°	38 <u>+</u> 3
16	1%	424 <u>+</u> 96°	1209±325°	31 <u>+</u> 5
24	1%	187 <u>+</u> 39	813 <u>+</u> 151 °	39 <u>+</u> 1
48	1%	141 <u>+</u> 24	145 <u>+</u> 48 °	36 <u>+</u> 6
			IFN-α	
0	untreated	<200	· NA	<650
16	placebo	<200	<650	<650
1	1%	<200	<650	<650
2	1%	<200	<650	<650
4	1%	<200	<650	<650
8	1%	<200	3/5>650	<650
16	1%	<200	<650	<650
24	1%	<200	<650	<650
48	1%	<200	<650	<650

^aFemale hairless CD rats were dosed topically with cream formulated Compound 1.

^bTNF-α and MCP-1 were measured by ELISA. IFN-α was measured by bioassay. Results are presented in pg/ml for serum samples and pg/200 mg tissue for tissue samples and represent the mean of five animals ± SEM.

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^cIndicates p<0.05 when compared to either placebo for serum samples or the difference between treated tissue and control tissue from the same animal.

A multiple dose study was done to monitor effects of a multiple dose regimen (Study 2 with results shown in Table 3). Rats were dosed two times a week for six hours for three weeks with topical cream formulation of Example 5. Placebo (Comparative Example C1) and single dosed rats were done for comparison and done simultaneously with the last dosing of the multiple dose set. Serum and tissue samples were taken at 8 and 24 hours post dose and analyzed for MCP-1.

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An analysis identical to that of Study 1 was performed for Study 2. This data set was broken up by treatment (multiple- or single-use) and time point prior to analysis. Again, placebo data were recorded only at the 8-hour time point for single use, but were used to compare placebo to every treatment and time point combination separately. The results are set forth in Table 3 below.

Table 3. Cytokine Concentrations in Rat Serum and Dermal Tissue Following Topical Application of the Topical Cream Formulation of Example 5 Multiple vs. Single Dose^a

	Dose	Cytokine Concentration ^b		
Time		MCP-1		
(hours)				
Post Dose	·	Serum	Treated Site	Control Site
0 .	None	89 <u>+</u> 11	NA	20 <u>+</u> 10
	(untreated)			_
24	Placebo	41 <u>+</u> 14	42 <u>+</u> 15	28 <u>+</u> 6
	Multiple			
8	0.1%	71 <u>+</u> 13	784 <u>+</u> 48 °	42 <u>+</u> 5
	Multiple]	
_ 24	0.1%	105 <u>+</u> 36	145 <u>+</u> 23 °	32 <u>+</u> 6
	Single			
8	0.1%	73 <u>+</u> 9	519 <u>+</u> 99°	33 <u>+</u> 6
· V	Single			
24	0.1%	82 <u>+3</u> °	412 <u>+</u> 130°	35 <u>+</u> 7

^aFemale hairless CD rats were dosed topically with cream formulated Compound 1. ^b MCP-1 was measured by ELISA. Results are presented in pg/ml for serum samples and pg/200 mg tissue for tissue samples and represent the mean of five animals \pm SEM. ^cIndicates p<0.05 when compared to either placebo for serum samples or the difference between treated tissue and control tissue from the same animal.

A dose response study (Study 3 with results shown in Table 4) was performed by dosing with the topical cream formulations of Examples 3-5 and 7, containing various concentrations of IRM Compound 1. Serum and tissue samples were taken at 8 and 24 hours post dose and analyzed for MCP-1. The studies tested topical delivery of creams comprising IRM Compound 1 for its ability to affect a local MCP-1 induction at four concentrations.

Serum data compared active treatment to placebo (Comparative Example C1) separately at each specified time point. Note that the placebo group was only measured at 24 hours post dose and these observations were compared to each time point for the active group.

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Table 4. Cytokine Concentrations in Rat Serum and Dermal Tissue Following Topical Application of the Formulations of Examples 3-5 and 7 an

		Cytokine Concentration ^b		
Time (hours)			MCP-1	
Post Dose	Dose	<u></u>		
		Serum	Treated Site	Control Site
0	controls	207 <u>+</u> 96	NA NA	38 <u>+</u> 12
24	placebo	367 <u>+</u> 178	61 <u>+</u> 14	20 <u>+</u> 5
	(Comparative			
	Example C1)			
8	0.01%	81 <u>+</u> 23	61 <u>+</u> 12	36 <u>+</u> 7
•	(Example 3)			
8	0.03%	81 <u>+</u> 20	· 271 <u>+</u> 29	48 <u>+</u> 5
	(Example 4)	•		
8	0.1%	153 <u>+</u> 14	1119±122°	51±8
	(Example 5)		1	
8	1.0%	136 <u>+</u> 23	1370 <u>+</u> 99°	50 <u>+</u> 15
	(Example 7)			
24	0.01%	71 <u>+</u> 18	183 <u>+</u> 49 °	33 <u>+</u> 13
	(Example 3)			
24	0.03%	71 <u>+</u> 20	212 <u>+</u> 49 °	40 <u>+</u> 7
	(Example 4)			
24	0.1%	226 <u>+</u> 73	628 <u>+</u> 127°	40 <u>+</u> 11
	(Example 5)			
24	1.0%	149 <u>+</u> 45	756 <u>+</u> 38 °	30 <u>+</u> 9
	(Example 7)			

^aFemale hairless CD rats were dosed topically with cream formulated Compound 1.

^b MCP-1 was measured by ELISA. Results are presented in pg/ml for serum samples and pg/200 mg tissue for tissue samples and represent the mean of five animals ± SEM.

^cIndicates p<0.05 when compared to either placebo for serum samples or the difference between treated tissue and control tissue from the same animal.

Examples 8-13

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Table 5 summarizes topical formulations made in accordance with the present invention in a percentage weight-by-weight basis.

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Ingredient	•		Topical Cream (percentage weight)	Cream ght-by-weight)		
(Compendial Status)	Example 8	Example 9	Example 10	Example 11	Example 12	Example 13
IRM Compound 2	0.01	0.03	0.10	1.00	0.003	0.30
Isostearic Acid	5.00	5.00	5.00	10.00	5.00	5.00
Isopropyl Myristate (NF)	10.00	10.00	10.00	5.00	10.00	10.00
Carbomer 974P (NF)	1.00	1.00	1.00	0.75	1.00	1.00
Poloxamer 188 (NF)	2.50	2.50	2.50	2.50	2.50	. 2.50
Propylene Glycol (USP)	15.00	15.00	15.00	15.00	15.00	15.00
Methylparaben (NF)	0.20	0.20	0.20	0.20	0.20	0.20
Propylparaben (NF)	0.10	0.10	0.10	0.10	0.10	0.10
Edetate Disodium (USP)	0.05	0.05	0.05	0.05	0.05	0.05
Sodium Hydroxide (NF) Solution, 20% w/w	0.50	0.50	0.50	0.35	05.0	0.50
Purified Water (USP)	65.64	65.62	65.55	65.05	65.647	:65.35
Total	100.00	100.00	100.00	100.00	100.00	100.00

The formulations set forth in Table 5 were prepared in the following manner:

Oil phase preparation: N-[4-(4-Amino-2-butyl-1*H*-imidazo[4,5-*c*] [1,5]naphthyridin-1-yl)butyl]-N'-cyclohexylurea (IRM Compound 2) was dissolved in isostearic acid and isopropyl myristate, with heat if necessary. Carbomer 974P was then dispersed in the oil phase.

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Water phase preparation: Edetate disodium was dissolved in the water.

Methylparaben and propylparaben were dissolved in propylene glycol, and the solution was subsequently added to the water phase. Poloxamer 188 was then added to the water phase and mixed until dissolved.

Phase combination: The oil phase was added to the water phase at ambient conditions. The emulsion was then homogenized. After homogenization, sodium hydroxide solution (20% w/w) was added and the resulting cream was mixed until smooth and uniform. The pH of the cream was measured, and a pH adjustment was made with additional sodium hydroxide solution, if necessary, to meet the in-process target pH of 5.

Formulations containing N-[4-(4-Amino-2-butyl-1H-imidazo[4,5-c][1,5]naphthyridin-1-yl)butyl]-N'-cyclohexylurea (IRM Compound 2) were tested for their ability to induce increases in cytokine concentrations in rats following topical application. This study was undertaken to evaluate cytokine induction following a single dosing of various strengths and timepoints or a multiple vs. single dosing of IRM Compound 2. The formulations described above were tested by examining tissue and serum concentrations of TNF- α , MCP-1 and IFN- α following drug treatment as described in Examples 1-7.

The data shown below in Tables 6-8 are from three separate experiments and analyzed to 1) measure pharmacokinetics by full time course, 2) measure dose response and 3) measure multiple vs. single dosing.

In order to determine the kinetics of local and systemic cytokine production following local administration of IRM Compound 2, the full time course study (Study 1 with results in Table 6) was done by topically dosing rats with the topical cream formulation of Example 11 as described in Examples 1-7. The data are presented in Table 6.

Table 6. Cytokine Concentrations in Rat Serum and Dermal Tissue Following Application of the Topical Formulation of Example 11 Full Time Course^a

		Cy	tokine Concentra	ation ^b
Time (hours)			TNF-α	
Post Dose	Dose	Serum	Treated Site	Control site
0	untreated	29+15	NA .	70+11
16	placebo	42+9	131 <u>+</u> 32	69+11
1	1%	38 <u>+</u> 38	44 <u>+</u> 14	35 <u>+</u> 19
2	1%	2+2	75±20°	33 <u>+</u> 13
4 .	1%	3 <u>+</u> 3	321 <u>+</u> 18°	62+20
8	1%	0	894 <u>+</u> 180°	21 <u>+</u> 9
16	1%	12 <u>+</u> 12	377 <u>+</u> 45°	22 <u>+</u> 12
24	1%	16 <u>+</u> 8	285±15°	52 <u>+</u> 14
48	1%	24 <u>+</u> 7	74 <u>+</u> 9	65±13
			MCP-1	•
0	untreated	100 <u>+</u> 20	NA	33 <u>+7</u>
16	placebo	144 <u>+</u> 9	225 <u>+</u> 106	22 <u>+</u> 4
1	1%	117 <u>+</u> 17	· 56 <u>+</u> 9	55 <u>+</u> 9
2	1%	126 <u>+</u> 29	50 <u>+</u> 13	54 <u>+</u> 8
4	1%	136±29	161±18°	71 <u>+</u> 9
8	1%	189 <u>+</u> 28	1020 <u>+</u> 319	45 <u>+</u> 15
16	1%	297 <u>+</u> 35	1294 <u>+</u> 122°	40 <u>+</u> 9
24	1%	217 <u>+</u> 12	1044 <u>+</u> 185 °	41 <u>+</u> 11
48	1%	120+22	134±14°	34 <u>+</u> 7
			IFN-α	. '
0	untreated	<65	· NA	<650
16	placebo	<65	<650	<650
1	1%	<65	<650	<650
2	1%	<65	<650	<650
4	1%	<65	<650	<650
8	1%	<65	901 <u>+</u> 571	<650
16	1%	<65	1330 <u>+</u> 386°	<650
24	1%	<65	<650	<650
48	1%	<65	<650	<650

^aFemale hairless CD rats were dosed topically with cream formulated Compound 2.

^bTNF-α and MCP-1 were measured by ELISA. IFN-α was measured by bioassay. Results are presented in pg/ml for serum samples and pg/200 mg tissue for tissue samples and represent the mean of five animals ± SEM.

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^cIndicates p<0.05 when compared to either placebo for serum samples or the difference between treated tissue and control tissue from the same animal.

A multiple dose study was done to monitor effects of a multiple dose regimen (Study 2 with results shown in Table 7). Rats were dosed two times a week for six hours for three weeks with topical cream formulation of Example 10. Placebo (Comparative Example C1) and single dosed rats were done for comparison and done simultaneously with the last dosing of the multiple dose set. Serum and tissue samples were taken at 16 and 24 hours post dose and analyzed for MCP-1.

An analysis identical to that of Study 1 was performed for Study 2. This data set was broken up by treatment (multi or single use) and time point prior to analysis. Again, placebo data were recorded only at the 16-hour time point for single use, but were used to compare placebo to every treatment and time point combination separately. The results are set forth in Table 7 below.

Table 7. Cytokine Concentrations in Rat Serum and Dermal Tissue Following Topical Application of the Topical Cream Formulation of Example 10 Multiple vs. Single Dose^a

		Cyt	okine Concentra	tion ^b
Time			MCP-1	
(hours) Post Dose	Dose	Serum	Treated Site	Control Site
0	None (untreated	161 <u>+</u> 58	NA	80 <u>+</u> 22
16	Placebo	214 <u>+</u> 35	71 <u>+</u> 16	47 <u>+</u> 11
16	Multiple 0.1%	321 <u>+</u> 62	1173 <u>+</u> 117°	86 <u>+</u> 14
24	Multiple 0.1%	217 <u>+</u> 43	388 <u>+</u> 80°	58 <u>+</u> 5
16	Single 0.1%	205 <u>+</u> 32	1448 <u>+</u> 241 °	77 <u>+</u> 15
24	Single 0.1%	279 <u>+</u> 45	1172 <u>+</u> 288°	90 <u>+</u> 15

^aFemale hairless CD rats were dosed topically with cream formulated Compound 2.

^b MCP-1 was measured by ELISA. Results are presented in pg/ml for serum samples and pg/200 mg tissue for tissue samples and represent the mean of five animals ± SEM.

^cIndicates p<0.05 when compared to either placebo for serum samples or the difference between treated tissue and control tissue from the same animal.

A dose response study (Study 3 with results shown in Table 8) was performed by dosing with the topical cream formulations of Examples 8-11, containing various concentrations of IRM Compound 2. Serum and tissue samples were taken at 16 and 24 hours post dose and analyzed for MCP-1. The studies tested topical delivery of creams comprising IRM Compound 2 for its ability to affect a local MCP-1 induction at four concentrations.

Serum data compared active treatment to placebo (Comparative Example C1) separately at each specified time point. Note that the placebo group was only measured at 16 hours post dose and these observations were compared to each time point for the active group.

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Table 8. Cytokine Concentrations in Rat Serum and Dermal Tissue Following Topical Application of the Formulations of Examples 8-11 a

		C	ytokine Concentrati	on ^b
Time (hours) Post Dose	Dose		MCP-1	
	Ī	Serum	Treated Site	Control Site
0	controls	293 <u>+</u> 23	NA	41 <u>+</u> 11
16	placebo (Comparative Example C1)	293 <u>+</u> 76	44 <u>+</u> 10	36 <u>+</u> 12
16	0.01% (Example 8)	276 <u>+</u> 50	257 <u>+</u> 85	57 <u>+</u> 20
16	0.03% (Example 9)	318 <u>+</u> 86	210 <u>+</u> 10	45 <u>+</u> 9
16	0.10% (Example 10)	529 <u>+</u> 141	2622 <u>+</u> 616 °	73 <u>+</u> 9
. 16	1.0% (Example 11)	345 <u>+</u> 51	3166 <u>+</u> 470°	71 <u>+</u> 11
. 24	0.01% (Example 8)	298 <u>+</u> 65	276 <u>+</u> 87	94 <u>+</u> 32
24	0.03% (Example 9)	253 <u>+</u> 34	427 <u>+</u> 238	28 <u>+</u> 14
24	0.10% (Example 10)	331 <u>+</u> 93	1461 <u>+</u> 264 °	19 <u>+</u> 7
24	1.0% (Example 11)	358 <u>+</u> 52	. 1952 <u>+</u> 185 °	17 <u>+</u> 6

^aFemale hairless CD rats were dosed topically with cream formulated Compound 2.

^b MCP-1 was measured by ELISA. Results are presented in pg/ml for serum samples and pg/200 mg tissue for tissue samples and represent the mean of five animals ± SEM.

^cIndicates p<0.05 when compared to either placebo for serum samples or the difference between treated tissue and control tissue from the same animal.

10 **Examples 14 – 18**

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Table 9 summarizes topical formulations made in accordance with the present invention on a percentage weight-by-weight basis.

Table 9

		Top	ical Cream	3				
Ingredients	(percentage weight-by-weight)							
	Ex. 14	Ex. 15	Ex. 16	Ex. 17	Ex. 18			
IRM Compound 1	0.01	0.10	1.00	3.00	1.00			
Isostearic Acid (874)	5.00	5.00	10.00	25.00	10.00			
*Diisopropyl dimer	10.00	10.00	5.00	5.00	<u>-</u> .			
dilinoleate					·			
**Caprylic/capric	-	•	-	_	5.00			
triglycerides			•					
Carbomer 980, NF	0.70	0.70	0.70	0.90	0.70			
Diethylene glycol	10.00	10.00	10.00	10.00	10.00			
monoethyl ether		*						
USA - NF								
Disodium EDTA, USP	0.05	0.05	0.05	0.05	0.05			
Poloxamer 188, NF	2.50	2.50	2.50	2.50	2.50			
Purified Water	70.94	70.85	69.95	52.55	69.95			
Methylparaben, NF	0.20	0.20	0.20	0.20	0.20			
Ethylparaben	0.20	0.20	0.20	0.20	0.20			
20% (w/w) NaOH	0.40	0.40	0.40	0.60	0.40			
Total % w/w	100.00	100.00	100.00	100.00	100.00			

^{*}Available under the trade name PRIPURE 3786 from Uniquema, New Castle, DE

5 <u>Examples 19 – 24</u>

Table 10 summarizes topical formulations made in accordance with the present invention on a percentage weight-by-weight basis.

^{**}Available under the trade name Crodamol GTCC-PN from Croda, Inc, Parsippany, NJ

Table 10

	Topical Creams							
Ingredients	(percentage weight-by-weight)							
	Ex. 19	Ex. 20	Ex. 21	Ex. 22	Ex. 23	Ex. 24		
IRM Compound 2	0.003	0.03	0.10	1.00	3.00	1.00		
Isostearic Acid (874)	5.00	5.00	5.00	10.00	25.00	10.00		
Diisopropyl dimer	10.00	10.00	10.00	5.00	5.00	-		
dilinoleate			'					
Caprylic/capric	- .	-	-	-	-	5.00		
triglycerides								
Carbomer 980, NF	0.70	0.70	0.70	0.70	0.60	0.70		
Diethylene glycol	10.00	10.00	10.00	10.00	10.00	10.00		
monoethyl ether		·		•				
USA - NF						•		
Disodium EDTA, USP	0.05	0.05	0.05	0.05	0.05	0.05		
Poloxamer 188, NF	2.50	2.50	2.50	2.50	2.50	2.50		
Purified Water	70.95	70.92	70.85	69.95	53.19	69.95		
Methylparaben, NF	0.20	0.20	0.20	0.20	0.20	0.20		
Ethylparaben	0.20	0.20	0.20	0.20	0.20	0.20		
20% (w/w) NaOH	0.40	0.40	0.40	0.40	0.26	0.40		
Total % w/w	100.00	100.00	100.00	100.00	100.00	100.00		

The formulations described in Tables 9 and 10 were prepared using the following general method:

Oil phase preparation:

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The IRM compound was dissolved in isostearic acid and diisopropyl dimer dilinoleate (or caprylic/capric acid triglyceride) with heat if necessary.

Water phase preparation:

Edetate disodium was dissolved in the water. Poloxamer 188 was then added to the water phase and mixed until dissolved. Carbomer 980 was then added to the water phase and mixed until the carbomer was fully dispersed and hydrated. Methylparaben and propylparaben were dissolved in diethylene glycol monoethyl ether and the solution was subsequently added to the water phase.

Phase combination:

The water phase was added to the oil phase at ambient conditions. The emulsion was then mixed at high speed or homogenized. After homogenization, sodium hydroxide solution (20% w/w) was added and the resulting cream was mixed until smooth and uniform. The pH of the cream was measured and a pH adjustment was made with additional sodium hydroxide solution, if necessary, to meet the in-process target pH of 5.

Examples 25 - 28

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Table 11 summarizes topical formulations made in accordance with the present invention on a percentage weight-by-weight basis.

Table 11

		Topica	al Cream				
Ingredient	(percentage weight-by-weight)						
	Ex. 25	Ex. 26	Ex. 27	Ex. 28			
IRM Compound 1	1	1	1	1			
Isostearic Acid (874)	10	10	10	8			
Diisopropyl dimer	5	5	5	1			
dilinoleate							
Carbomer 980, NF	0.7	0.7	0.7	0.7			
Diethylene glycol	10	10	10	10			
monoethyl ether							
USA - NF							
Disodium EDTA, USP	0.05	0.05	0.05	0.05			
Poloxamer 188, NF	2.5	2.5	2.5	2.5			
Purified Water	Qs to 100	Qs to 100	Qs to 100	Qs to 100			
Methylparaben, NF	0.2	0.2	0.2	0.2			
Ethylparaben	0.2	0.2	0.2	0.2			
20% (w/w) NaOH	0.4	0.4	0.4	0.4			
10% iodopropynyl	-	.1	+	- ;			
butylcarbamate in							
PEG-4 laurate	. •						
Phenoxyethanol	-	-	0.5	-			

Examples 29 - 135

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Topical creams containing the IRM compounds listed in Table 12 were prepared using the general methods described above for Examples 1-24. Each IRM was formulated into one or more of the model formulations shown in Tables 13 and 14. Table 15 summarizes the topical creams that were prepared.

Table 12

IRM	Chemical Name
Compound	·
3	1-(2-methylpropyl)-1H-imidazo[4,5-c]quinolin-4-amine
4	1-(2-methylpropyl)-1 <i>H</i> -imidazo[4,5- <i>c</i>][1,8]naphthyridin-4-amine
5	2-butyl-1-(2-methylpropyl)-1 <i>H</i> -imidazo[4,5-c][1,8]naphthyridin-4-amine
6	1-(2-methylpropyl)-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine
7 ·	2-methylthiazolo[4,5-c]quinolin-4-amine
8	2-ethoxymethyl-1-phenylmethyl-1H-imidazo[4,5-c][1,5]naphthyridin-4-
	amine
9	2-ethylthiazolo[4,5-c]quinolin-4-amine
10.	4-amino-2-butyl- α , α -dimethyl-1 H -imidazo[4,5- c][1,5]naphthyridine-1-
	ethanol
11	N ¹ -[2-(4-amino-1 <i>H</i> -imidazo[4,5-c]quinolin-1-yl)ethyl]benzamide
12	1-{2-[3-(3-pyridyl)propoxy]ethyl}-1H-imidazo[4,5-c]quinolin-4-amine
13	1-(2-phenoxyethyl)-1 <i>H</i> -imidazo[4,5- <i>c</i>]quinolin-4-amine
14	1-[(R)-1-phenylethyl]-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine
15	N^4 -[4-(4-amino-1 <i>H</i> -imidazo[4,5- <i>c</i>]quinolin-1-yl)butyl]-4-
	morpholinecarboxamide
16	N ³ -[4-(4-amino-1 <i>H</i> -imidazo[4,5- <i>c</i>]quinolin-1-yl)butyl]nicotinamide
17	$1-\{2-[3-(1,3-\text{thiazol-}2-\text{yl})\text{propoxy}]\text{ethyl}\}-1H-\text{imidazo}[4,5-c]\text{quinolin-}4-\text{amine}$
18	1-[2-(pyridin-4-ylmethoxy)ethyl]-1 <i>H</i> -imidazo[4,5- <i>c</i>]quinolin-4-amine
19	2-methyl-1-[5-(methylsulfonyl)pentyl]-1H-imidazo[4,5-c]quinolin-4-amine
20	N-[3-(4-amino-2-methyl-1 <i>H</i> -imidazo[4,5- <i>c</i>]quinolin-1-yl)propyl]
	cyclohexanecarboxamide
21	N-[3-(4-amino-2-methyl-1 <i>H</i> -imidazo[4,5- <i>c</i>]quinolin-1-yl)propyl]-2-
	methylpropanamide
22	N-[3-(4-amino-2-methyl-1 <i>H</i> -imidazo[4,5-c]quinolin-1-yl)propyl]butanamide

IRM	Chemical Name
Compound	·
23	2-butyl-1-{2-[(1-methylethyl)sulfonyl]ethyl}-1H-imidazo[4,5-c]quinolin-4-
	amine
24	N-{2-[4-amino-2-(ethoxymethyl)-1 <i>H</i> -imidazo[4,5-c]quinolin-1-yl]ethyl}
	ethanesulfonamide
25	N-{2-[4-amino-2-(ethoxymethyl)-1H-imidazo[4,5-c]quinolin-1-yl]ethyl}
	propanamide
26	1-[2-(methylsulfonyl)ethyl]-2-propyl-1 <i>H</i> -imidazo[4,5- <i>c</i>]quinolin-4-amine
27	N-{2-[4-amino-2-(ethoxymethyl)-1H-imidazo[4,5-c]quinolin-1-yl]ethyl}-N'-
	ethylthiourea
28	2-ethyl-1-{4-[(1-methylethyl)sulfonyl]butyl}-1H-imidazo[4,5-c]quinolin-4-
	amine
29	2-ethyl-1-[4-(ethylsulfonyl)butyl]-1 <i>H</i> -imidazo[4,5- <i>c</i>]quinolin-4-amine
30	$N-\{3-[4-amino-2-(ethoxymethyl)-1H-imidazo[4,5-c]quinolin-1-yl]propyl\}$
	cyclopentanecarboxamide
31	N-{3-[4-amino-2-(ethoxymethyl)-6,7-dimethyl-1 <i>H</i> -imidazo[4,5-c]pyridin-1-
	yl]propyl}morpholine-4-carboxamide
32	1-(2-methylpropyl)-6,7,8,9-tetrahydro-1 <i>H</i> -imidazo[4,5- <i>c</i>]quinolin-4-amine
33	8,9,10,11-tetrahydropyrido[1',2':1,2]imidazo[4,5-c]quinolin-6-amine
34	4-amino- α , α , 2-trimethyl-6, 7, 8, 9-tetrahydro-1 H -imidazo[4, 5- c] quinoline-1-
	ethanol
35	2-hydroxymethyl-1-(2-methylpropyl)-6,7,8,9-tetrahydro-
	1 <i>H</i> -imidazo[4,5-c]quinolin-4-amine
36	2-butyl-1-(2-phenoxyethyl)-1 <i>H</i> -imidazo[4,5-c][1,5]naphthyridin-4-amine
37	N-[3-(4-amino-2-methyl-1 <i>H</i> -imidazo[4,5- <i>c</i>]quinolin-1-yl)propyl]
	methanesulfonamide
L	

Table 13

	Model Formulation								
Ingredient		(percentage weight-by-weight)							
	A	В	С	D	Е	F	G		
IRM	0.01	0.1	1	1	1	1	1		
Isostearic acid	5	5	5	20	42	13	6		
Isopropyl myristate	10	10	10	10	2	10	10		
Carbomer 974P	1	1	1	1	1	1.5	1		
Purified water	*	*	*	*	*	*	*		
Poloxamer 188	2.5	2.5	2.5	2.5	2.5	2.5	2.5		
Propylene glycol	15	15	15	15	1:3	15	15		
Xanthan gum	-	-	-	-	0.4	•	-		
Methylparaben	0.2	0.2	0.2	0.2	0.2	0.2	0.2		
Disodium EDTA	0.05	0.05	0.05	0.05	0.05	0.05	0.05		
20% NaOH	0.7	0.7	0.7	0.7	0.7	0.7	0.7		

^{*}Qs to 100

Table 14

		Mo	odel For	mulatio	n	
Ingredient	(percent	age wei	ght-by-	weight)	
	Н	I	J	K	L	M
IRM	0.01	0.1	1	1	3	5
Isostearic acid	5	5	5	10	. 10	10
Diisopropyl dimer	10	10	10	- 5	5	5
dilinoleate						
Carbomer 980	0.7	0.7	0.7	1.0	1.0	1.0
Purified water	*	*	*	*	*	*
Poloxamer 188	2.5	2.5	2.5	2.6	2.6	2.6
Diethylene glycol	10	10	10	10	10	10
monoethyl ether				•		
Xanthan gum	-	-	-	0.1	0.1	0.1
Methylparaben	0.2	0.2	0.2	0.2	0.2	0.2
Ethylparaben	0.2	0.2	0.2	0.2	0.2	0.2
Disodium EDTA	0.05	0.05	0.05	0.05	0.05	0.05
20% NaOH	0.4	0.4	0.4	0.4	0.4	0.4

^{*}Qs to 100

Table 15

Example	IRM Compound	Model Formulation
29	3	A
30	3	В
31	3	C.
32	4	A
33	. 4	В
34	4	C
35	5	A

Example	IRM Compound	Model Formulation
36	5	В
37	5	D
38	6	A
39	6	В
40	6	С
41	7	A
42	7	В
43	7	С
44	8	A
45	8	· B
46	8	С
47	9	A
48	9	В
49	9	С
.50	10	A
51	10	В
52	10	С
53 .	11 ,	A
54	11	В
55	11	Е
56	12	Α .
57	12	В
58	12	С
59	13	A
60	13	В
61	13	F
62	14	A
63	. 14	В

Example	IRM Compound	Model Formulation
64	14	G
65	15	Н
66	15	I
67	15	K
68	16	Н
69	16	I
70	16	K
71	17	A
72	17	В
73	17	· C
74	18	Н
75	. 18	I
76	18	K
77	19	Н
78	19	I
79	19	K
80	20	Н
81	20 .	. I
82	20	К
83	20	L
84	20	M
85	21	Н
86	.21	. I
87	21	K
88	22	H
89	22	I
90	22	J
91	23	Н

Example	IRM Compound	Model Formulation
92	23	I
93	23	J
94	24	Н
95	24	I
96	24	. К
97	25.	Н
98	25	I
99.	25	K
100	26	H
101	26	. I
102	26	K
103	27	Н
104	27	I
105	27 .	K
106	28	Н
107	28	I
108	28	K
109	29 ,	Н
110	29	I
111	29	K
112	30	Н
113	30	I
114	30	K
115	31	Н
116	31	. I
117	31	К
118	32	A
119	, 32	В

Example	IRM Compound	Model Formulation
120	32	C
121	33	A
122	33	В
123	33	C
124	34	A
125	34	В
126	34	, C
127	35	A
128	35	В
129	35	С
130	36	A
131	36	В
132	36	С
133	37	Н
· 134	. 37	I
135	37	K

The topical creams of Examples 29 –135 were tested using the test method described below. The results are shown in Table 16 below where each value is the mean of the values from the 3 rats in the treatment group.

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SINGLE DOSE MCP-1 INDUCTION TEST METHOD

Female CD hairless rats (Charles River Laboratories, Wilmington, MA) weighing 200-250 grams are used. Animals are randomized to treatment groups and dosed three per treatment group.

The rats are acclimated to collars around the neck on two consecutive days prior to actual dosing. A 50 μ L dose of active cream or the appropriate placebo is applied to the right flank and gently rubbed into the skin of the rat. The rats are then collared and housed

individually to prevent ingestion of the drug. At selected post treatment time points, the rats are anesthetized, and blood (3 mls) is collected by cardiac puncture. Blood is allowed to clot at room temperature. Serum is separated from the clot via centrifugation, and stored at -20°C until it is analyzed for MCP-1 concentration.

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Following blood collection, the rats are euthanized, and their skins removed. Tissue samples (4 from each site) from both the treated site and contralateral site (untreated) are obtained using an 8 mm punch biopsy, weighed, placed in a sealed 1.8 ml cryovial, and flash frozen in liquid nitrogen. The frozen tissue sample is then suspended in 1.0 mL RPMI medium (Celox, Hopkins, MN) containing 10% fetal bovine serum (Sigma, St. Louis, MO), 2 mM L-glutamine, penicillin/streptomycin, and 2-mercaptoethanol (RPMI complete) combined with a protease inhibitor cocktail set III (Calbiochem, San Diego, CA). The tissue is homogenized using a Tissue TearorTM (Biospec Products, Bartlesville, OK) for approximately 1 minute. The tissue suspension is then centrifuged at 2000 rpm for 10 minutes under refrigeration to pellet debris, and the supernatant is collected and stored at -20°C until analyzed for MCP-1 concentration.

ELISAs for rat MCP-1 are purchased from BioSource Intl. (Camarillo, CA) and performed according to manufacturer's specifications. Results are expressed in pg/ml, the values for the tissue samples are normalized per 200 mg of tissue. The sensitivity of the MCP-1 ELISA is 12 pg/ml.

able 16

						<u> </u>			r				
	Cream		Untreated	59	65	59	56	. 95	99	28	58	58	59
	Placebo Cream	:`	Serum	142	142	142	54	54	54	08	80	80	142
			Untreated	34	43	125	69	58	69	62	96	77	98
		24 hours	Treated	55	201	909	. 28	74	319	142	1216	2036	121
MCP-1 (pg/ml)			Serum	291	177	267	97	56	59	162	98	136	211
2	IRM Cream		Untreated	46	31	. 54	59	71	58	55	46	38	120
		6 hours	Treated	202	92	1235	54	70	88	110	674	1826	59
			Serum	123	119	212	26	54	72	170	94	153	178
		Cream of	Example	29	30	31	32	33	34	35	36	37	38

				,		r	r					·		Γ	
	Cream		Untreated	59	59	96	96	96	**28	**28	**28	73	73	73	*
	Placebo Cream	٠	Serum	142	142	73	73	73	128	128	128	37	37	37	88
			Untreated	59	95	71	88	93	*	*	*	73	61	75	56
		24 hours	Treated	263	1086	888	126	1016	488	1041	1023	95	112	1436	655
MCP-1 (pg/ml)			Serum	259	284	45	54	68	177	157	406	81	.63	83	*
Z	IRM Cream		Untreated	61	58	96	78	89 .	*	#	*	59	61	62	56
		6 hours	Treated	220	1204	82	129	824	256	1444	1720	53	200	1254	1033
	•		Serum	193	226	54	65	77	98	172	177	58	7.1	92	170
		Cream of	Example	39	. 40	41	42	43	44	45	46	47	48	49	50

				·							r				
	Placebo Cream		Untreated	*	*	31	31	31	**42	**42	**42	54	54	54	26
	Placebo		Serum	888	88	7	۲.	7	201	201	201	107	107	107	96
			Untreated	787	314	45	24	32	#	*	*	53	69	46	. 61
		24 hours	Treated	149	98	47	26	33	425	1252	1508 ·	41	80	1131	428
MCP-1 (pg/ml)			Serum	*	*	76	71	44	.115	267	476	271	175	151	174
2	IRM Cream		Untreated	787	314	46	27	. 21	*	*	*	36	58 .	52	55
		6 hours	Treated	551	348	63	35	35	44	411	1560	46.	53.	172	211
			Serum	625	811	70	89	75	115	119	190	155	. 123	133	143
		Cream of	Example	51	- 52	53	54	55	56	57	58	59	09	61	62

	Cream		Untreated	26		. 59	. 59	59	72	72	72	. 53	53	53	91
	Placebo Cream		Serum	96	. 96	83	83	83	89	89	. 89	177	177	177	61
-			Untreated	74	66	61	74	72	35	59	09	57	63	108	88
		24 hours	Treated	1217	390	81	. 42	34	72	73	134	84	066	1411	66
MCP-1 (pg/ml)			Serum	230	425	46	32	25	91	66	91	134	122	. 293	73
X	IRM Cream		Untreated	51	529	57	28	. 61	82	52	34	62	65	63	7.1
		6 hours	Treated	1614	1094	34	73	54	77	74		79	255	. 666	99
			Serum	320	970	43	29	19	09	143	59	259	138	251	66
	-	Cream of	Example	63	- 64	65	. 99		89	69	70	71	72	73	74

	Cream		Untreated	91	91	38	38	38	15	15	15	120	120	120	0
-	Placebo Cream		Serum.	61	.01	30	30	30	76			93	93	93	177
			Untreated	73	104	40	59	38	0	10	45	92	172	40	0
		24 hours	Treated	170	4949	43	49	50	163	34	303	225	275	629	0
MCP-1 (pg/ml)	*		. Serum	m .	188	21	33	27	50	83	121	79	61	86	0
M	IRM Cream		Untreated	78	. 64	35	37	. 40	23	15	0	36	124	. 92	0 :
		6 hours	Treated	101	6779	47	35	41	59	0	32	149	164	177	0
٠			Serum	94.	99	28	27	24	51	6	61	50	110	. 65	81
,		Cream of	Example	75	- 76	77	78	. 62	08	81	82***	82***	83.	84	85

						,									<u> </u>
	Placebo Cream		Untreated	0	0	33	. 33	33 .	46	46	46	. 100	100	. 100	102
	Placebo		Serum	177	177	141	141	141	34	34	34	110	110	110	36
			Untreated	0	0	42	40	41	57	29	74	133	112	81	92
		24 hours	Treated	0	0	43	49	109	874	1087	1124	136	158	528	95
MCP-1 (pg/ml)			Serum	0	0	87	132	111	86	92	114	48	95	132	54
A	IRM Cream		Untreated	0	0	41	49	. 47	53	74	64	. 107	06	79	. 106
		6 hours	Treated	0	0	56	47	96	91.	1238	2037	86	130	255	. 88
-			Serum	116	69	114	74	91	42	83	. 86	102	49	89	34
		Cream of	Example	98	- 87	88	68	06	91	92	93 .	94	56.	96	16

			g	1											7		
	Cream		Untreated	-	701	102	58	58	28	61	61	61	137	137	137	. 31	
-	Placebo Cream		Serum		36	36	82	82	82	7	7	7	∞	∞ ·	∞	12	
			Tintreated	Omnomic	91	9/	72	64	103	94	63	69	158	75	128	62	
		24 hours	Trooted	Tealen	123	945	115	209	3199	107	72	. 09	168	168	3267	78	
MCP-1 (pg/ml)				Serum	83	43	55	75	112	31	61	54	64	48	135	71	
Z	IRM Cream		•	Untreated	108	68	83	55	. 54	69	55	87	103	86	72	. 71	
		6 hours	-	Treated	116	150	81	72	489	88	. 99	83	96	129	314	09	
				Serum	17	51	111	33	79	. 82	13	75	72	21	95	72	
		Jo mood	Cicam or	Example	86	66 -	100	101	102	103	104	105	106	. 107	108	109	

	Cream	• .	Untreated	31	31	102	102	102	157	157	157	*	*	*	57
-	Placebo Cream		Serum	12	12	30	30	30	84	84	84	*	*	*	78
			Untreated	75	89	70	93	111	06	223	61	52	. 42	56	75
		24 hours	Treated	72	2397	84	1034	2880	77	57	91	46	104	892	96
MCP-1 (pg/ml)			Serum	92	32	28	70	196	84	64	62	135	144	171	888
X	RM Cream		Untreated	57	, 83	120	106	59	41	06	52	. 65	31	45	. 51
		6 hours	Treated	. 92	143	29	107	627	38	81	113	- 65	184	1261	74
	-		Serum	44	70	99	46	14	39	73	99	132	123	124	06
		Cream of	Example	110	- 111	112	113	114	115	116	. 117	118	119	120	. 121

						,		·	, ——,						T	1
	Cream		Untreated	57	. 57	652	652	652	. 81	81	81	32	32	. 32	∞	
	Placebo Cream		Serum	78	. 78	- 24	- 62	97	51	51	51	74	74	74	. 26	
			Untreated	82	. 48	110	120	349	22	35	59	43	09	11	32	
		24 hours	Treated	613	1043	95	72	1348	06	53	. 608	162 .	822	1212	199	
MCP-1 (pg/ml)			Serum	91	. 226	96	107	73	81	55	35	135	144	171	52	
Σ.	IRM Cream		Untreated	50	52	27	128	. 97	46	58	58	41	59	13	0	
		6 hours	Treated	415	1502	94	198	1828	99	80	382	55	279	1901	106	
			Serum	72	156	92	123	136		63	49	132	124	124	64	
		Cream of	Example	122	. 123	124	125	126	127	128	129	130	131	132	133	

MCP-1 (pg/ml)	Placebo Cream	IRM Cream	6 hours 24 hours		8 26 8	>	77 0 . 26 8)	
		IRM Cream	6 hours	-		· 		0 68	
	-	- 180		Serum		6		59	
			Cream of	Example		134		- 135	_

*MCP-1 concentration was not measured

**MCP-1 concentration is for the treated site.

***The cream of Example 82 was used in 2 separate experiments

Examples 136 - 140

Table 17 summarizes topical formulations made in accordance with the present invention on a percentage weight-by-weight basis.

Table 17

	Topical Creams (percentage weight-by-weight)							
Ingredients								
	Ex. 136	Ex. 137	Ex. 138	Ex. 139	Ex. 140			
IRM Compound 1	1	1	1	1	1			
Isostearic Acid	10	10	8 .	10	10			
Diisopropyl dimer		5	. 1	.5	5			
dilinoleate								
Caprylic/capric	5	-	-	-	-			
triglycerides								
Carbomer 980	0.7	0.7	0.7	0.7	0.7			
Diethylene glycol	10	10	10	10	10			
monoethyl ether				4				
Disodium EDTA	0.05	0.05	0.05	0.05	0.05			
Poloxamer 188	2.5	2.5	2.5	2.5	2.5			
Purified Water	Qs to 100	Qs to 100	Qs to 100	Qs to 100	Qs to 100			
Methylparaben	0.2	0.1	0.2	0.2	0.2			
Ethylparaben	0.2	0.1	0.2	0.2	0.2			
20% (w/w) NaOH	Qs to pH	Qs to pH	Qs to pH	Qs to pH	Qs to pH			
	5-5.5	5 – 5.5	5 – 5.5	6.5	5 – 5.5			
Iodopropynyl	-	0.1	-	-	-			
butylcarbamate	-							
PEG-4 Laurate	-	0.9	-	-	-			
Phenoxyethanol	-	1	-	: -	-			
Sorbic acid		0.15	-	-				

The topical creams of Examples 136 –140 were tested using the test method described below. The results are shown in Table 18 below where each value is the mean of the values from the 3 rats in the treatment group. "Normal animals" did not receive any treatment.

SINGLE DOSE CYTOKINE INDUCTION TEST METHOD

Female CD hairless rats (Charles River Laboratories, Wilmington, MA) weighing 200-250 grams are used. Animals are randomized to treatment groups and dosed three per treatment group.

The rats are acclimated to collars around the neck on two consecutive days prior to actual dosing. A 50 μ L dose of active cream is applied to the right flank and gently rubbed into the skin of the rat. The rats are then collared and housed individually to prevent ingestion of the drug. At 6 hours post treatment, the rats are anesthetized, and blood (3 mls) is collected by cardiac puncture. Blood is allowed to clot at room temperature, serum is separated from the clot via centrifugation, and stored at

-20°C until it is analyzed for cytokine concentrations.

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Following blood collection, the rats are euthanized, and their skins removed. Tissue samples (4 from each site) from both the treated site and contralateral site (untreated) are obtained using an 8 mm punch biopsy, weighed, placed in a sealed 1.8 ml cryovial, and flash frozen in liquid nitrogen. The frozen tissue sample is then suspended in 1.0 mL RPMI medium (Celox, Hopkins, MN) containing 10% fetal bovine serum (Sigma, St. Louis, MO), 2 mM L-glutamine, penicillin/streptomycin, and 2-mercaptoethanol (RPMI complete) combined with a protease inhibitor cocktail set III (Calbiochem, San Diego, CA). The tissue is homogenized using a Tissue TearorTM (Biospec Products, Bartlesville, OK) for approximately 1 minute. The tissue suspension is then centrifuged at 2000 rpm for 10 minutes under refrigeration to pellet debris. The supernatant is collected and stored at -20°C until analyzed for cytokine concentrations.

ELISAs for rat MCP-1 are purchased from BioSource Intl. (Camarillo, CA) and rat TNF-α are purchased from BD Pharmingen (San Diego, CA) and performed according to

manufacturer's specifications. Results are expressed in pg/ml, the values for the tissue samples are normalized per 200 mg of tissue. The sensitivity of the MCP-1 ELISA is 12 pg/ml and the sensitivity of the TNF- α ELISA is 31 pg/ml.

Table 18

1		TNF-a (pg/ml)	Tissue		29		29	67
	Normal Animals		Serum	49	64	64	64	64
	Normal	MCP-1 (pg/ml)	Tissue	39	39	39	39	39
		MCP-	Serum	73	73	73	73	73
Cytokine Induction		ml)	Untreated	85	78	69	75	95
		TNF-α (pg/ml)	Serum Treated	808	597	636	443	948
	Animals		Serum	64	78	99	50	80
	RM Cream Treated Animals	(III)	Untreated	5.1	78	27	. \$8	28
	IRM Cr	MCP-1 (pg/ml)	Treated	1208.	1815	1351	1509	2373
		V.	Serum	119	06	5	.62	24
		Cream of	Example	- 136	137	138	139	140

What is Claimed is:

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1. A pharmaceutical formulation comprising: an immune response modifier (IRM) compound selected from imidazoquinoline amines, imidazotetrahydroquinoline amines, imidazopyridine amines, 6,7-fused

cycloalkylimidazopyridine amines, 1,2- bridged imidazoquinoline amines, thiazoloquinoline amines, oxazoloquinoline amines, thiazolopyridine amines, oxazolopyridine amines, imidazonaphthyridine amines, imidazotetrahydronaphthyridine amines, and thiazolonaphthyridine amines;

a fatty acid;

a hydrophobic, aprotic component miscible with the fatty acid and comprising a hydrocarbyl group of 7 or more carbon atoms; and

a hydrophilic viscosity enhancing agent selected from cellulose ethers and carbomers.

- 2. The formulation according to claim 1 wherein the formulation further comprises a preservative system and an emulsifier.
 - 3. The formulation according to claim 1 wherein the hydrophobic, aprotic component has a hydrophilic lipophilic balance of less than 2.
- 4. The formulation according to claim 1 wherein the hydrophobic, aprotic component has a pKa greater than 14.2.
 - 5. The formulation according to claim 1 wherein the ratio of the hydrophobic, aprotic component to the fatty acid is 0.025:1 to 600:1.
 - 6. The formulation according to claim 1 wherein the combined weight percent of the hydrophobic, aprotic component and the fatty acid is 2 to 50.
 - 7. The formulation according to claim 1 wherein the fatty acid is isostearic acid.

8. The formulation according to claim 1 wherein the hydrophobic, aprotic component is selected from aprotic fatty acid esters, hydrocarbons of 8 or more carbon atoms, and waxes.

- 5 9. The formulation according to claim 8 wherein the aprotic fatty acid ester is isopropyl myristate, isopropyl palmitate, diisopropyl dimer dilinoleate, caprylic/capric triglyceride, cetyl esters wax, or a combination thereof.
- 10. The formulation according to claim 8 wherein the hydrocarbon of 8 or more carbon atoms is mineral oil or petrolatum.
 - 11. The formulation according to claim 1 wherein the hydrophilic viscosity enhancing agent comprises a carbomer.
- 15 12. The formulation according to claim 2 wherein the preservative system comprises methylparaben at 0.01 to 0.5% w/w of the formulation and propylparaben at 0.01 to 0.5% w/w of the formulation.
- 13. The formulation according to claim 2 wherein the preservative system comprises methylparaben at 0.01 to 0.5% w/w of the formulation and ethylparaben at 0.01 to 0.5% w/w of the formulation.

- 14. The formulation according to claim 2 wherein the preservative system comprises iodopropynyl butylcarbamate.
- 15. The formulation according to claim 2 wherein the preservative system comprises iodopropynyl butylcarbamate and one or more of methylparaben, ethylparaben, propylparaben, or phenoxyethanol.
- 30 16. The formulation according to claim 2 wherein the preservative system

comprises iodopropynyl butylcarbamate, methylparaben, and ethylparaben.

17. The formulation according to claim 2 wherein the preservative system comprises phenoxyethanol and one or both of methylparaben and ethylparaben.

18. The formulation according to claim 2 wherein the preservative system comprises a preservative enhancing solubilizer.

The formulation according to claim 18 wherein the preservative enhancing
 solubilizer comprises diethylene glycol monoethyl ether, propylene glycol or a combination thereof.

20. The formulation of claim 2 comprising:

- (a) 0.001 to 5% w/w
- 2-methyl-1-(2-methylpropyl)-1*H*-imidazo[4,5-*c*] [1,5]naphthyridin-4-amine,
 - N-[4-(4-amino-2-butyl-1H-imidazo[4,5-c][1,5]naphthyridin-1-yl)butyl]-N'-cyclohexylurea,
 - 1-(2-methylpropyl)-1H-imidazo[4,5-c]quinolin-4-amine,
 - 2-butyl-1-(2-methylpropyl)-1H-imidazo[4,5-c][1,8]naphthyridin-4-amine,
 - 1-(2-methylpropyl)-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
- 20 2-methylthiazolo[4,5-c]quinolin-4-amine,
 - 2-ethoxymethyl-1-phenylmethyl-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
 - 2-ethylthiazolo[4,5-c]quinolin-4-amine,
 - 4-amino-2-butyl- α , α -dimethyl-1*H*-imidazo[4,5-c][1,5]naphthyridine-1-ethanol,
 - 1-{2-[3-(3-pyridyl)propoxy]ethyl}-1H-imidazo[4,5-c]quinolin-4-amine,
- 25 1-(2-phenoxyethyl)-1*H*-imidazo[4,5-c]quinolin-4-amine,
 - 1-[(R)-1-phenylethyl]-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
 - $1-\{2-[3-(1,3-\text{thiazol-}2-yl)\text{propoxy}]\text{ethyl}\}-1H-\text{imidazo}[4,5-c]\text{quinolin-}4-\text{amine},$
 - 1-[2-(pyridin-4-ylmethoxy)ethyl]-1H-imidazo[4,5-c]quinolin-4-amine,

 $\label{eq:N-solution} $$N-[3-(4-amino-2-methyl-1$H-imidazo[4,5-c]$ quinolin-1-yl) propyl]$$ cyclohexanecarboxamide,$

 $2-\text{butyl-1-}\{2-[(1-\text{methylethyl})\text{sulfonyl}] \text{ethyl}\}-1\\ H-\text{imidazo}[4,5-c] \text{quinolin-4-amine,}$

 $N-\{2-[4-amino-2-(ethoxymethyl)-1 \\ H-imidazo[4,5-c] \\ quinolin-1-yl] \\ ethyl\}$

5 ethanesulfonamide,

N- $\{2-[4-amino-2-(ethoxymethyl)-1H-imidazo[4,5-c]quinolin-1-yl]ethyl\}$ propanamide,

1-[2-(methylsulfonyl)ethyl]-2-propyl-1H-imidazo[4,5-c]quinolin-4-amine,

2-ethyl-1-{4-[(1-methylethyl)sulfonyl]butyl}-1H-imidazo[4,5-c]quinolin-4-amine,

2-ethyl-1-[4-(ethylsulfonyl)butyl]-1*H*-imidazo[4,5-*c*]quinolin-4-amine,
N-{3-[4-amino-2-(ethoxymethyl)-1*H*-imidazo[4,5-*c*]quinolin-1-yl]propyl}

cyclopentanecarboxamide,

 $1-(2-\mathrm{methylpropyl})-6,7,8,9-\mathrm{tetrahydro-}1H-\mathrm{imidazo}[4,5-c]\mathrm{quinolin-}4-\mathrm{amine},$

8,9,10,11-tetrahydropyrido[1',2':1,2]imidazo[4,5-c]quinolin-6-amine,

4-amino- α , α , 2-trimethyl-6, 7, 8, 9-tetrahydro-1H-imidazo[4, 5-c] quinoline-1-ethanol,

2-hydroxymethyl-1-(2-methylpropyl)-6,7,8,9-tetrahydro-

1H-imidazo[4,5-c]quinolin-4-amine,

2-butyl-1-(2-phenoxyethyl)-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine, or a combination thereof;

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- (b) 0.05 to 40% w/w isostearic acid;
- (c) 1 to 30% w/w hydrophobic, aprotic component;
- (d) 0.5 to 10% w/w emulsifier;
- (e) 0.01 to 30% w/w preservative system; and
- (f) 0.1 to 10% carbomer.

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- 21. The formulation of claim 20 comprising:
- (a) 0.03 to 3% w/w 2-methyl-1-(2-methylpropyl)-1*H*-imidazo[4,5-*c*] [1,5]naphthyridin-4-amine, N-[4-(4-amino-2-butyl-1*H*-imidazo[4,5-*c*][1,5]naphthyridin-1-yl)butyl]-N'-cyclohexylurea, 2-butyl-1-{2-[(1-methylethyl)sulfonyl]ethyl}-1*H*-imidazo[4,5-*c*]quinolin-4-amine,or a combination thereof;

- (b) 3 to 25% w/w isostearic acid;
- (c) 3 to 15% w/w hydrophobic, aprotic component;
- (d) 0.75 to 3.5% w/w emulsifier;
- (e) 0.1 to 25% w/w preservative system; and
- (f) 0.5 to 5% w/w carbomer.

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- 22. A method of treating a dermal associated condition, the method comprising a step of: applying to skin a formulation comprising an immune response modifier (IRM) selected from imidazoquinoline amines, imidazotetrahydroquinoline amines, imidazopyridine amines, 6,7-fused cycloalkylimidazopyridine amines, 1,2-bridged imidazoquinoline amines, thiazoloquinoline amines, oxazoloquinoline amines, thiazolopyridine amines, oxazolopyridine amines, imidazonaphthyridine amines, imidazotetrahydronaphthyridine amines, and thiazolonaphthyridine amines; a fatty acid; a hydrophobic, aprotic component miscible with the fatty acid and comprising a hydrocarbyl group of 7 or more carbon atoms; and a hydrophilic viscosity enhancing agent selected from cellulose ethers and carbomers.
 - 23. The method according to claim 22 wherein the ratio of the hydrophobic, aprotic component to the fatty acid is 0.025:1 to 600:1.
 - 24. The method according to claim 22 wherein the combined weight percent of the hydrophobic, aprotic component and the fatty acid is 2 to 50.
- 25. The method according to claim 22 wherein the hydrophobic, aprotic component is selected from the group consisting of aprotic fatty acid esters, hydrocarbons of 8 or more carbon atoms, and waxes.
 - 26. The method according to claim 25 wherein the aprotic fatty acid ester is isopropyl myristate, isopropyl palmitate, diisopropyl dimer dilinoleate, caprylic/capric triglyceride, cetyl esters wax, or combinations thereof.

- 27. The method according to claim 22 wherein the hydrophilic viscosity enhancing agent comprises a carbomer.
- The method according to claim 22 wherein the topical formulation further comprises:

a preservative system; and an emulsifier.

- The method according to claim 22 wherein the IRM is 2-methyl-1-(2-methylpropyl)-1H-imidazo[4,5-c] [1,5]naphthyridin-4-amine, N-[4-(4-amino-2-butyl-1H-imidazo[4,5-c][1,5]naphthyridin-1-yl)butyl]-N'-cyclohexylurea, 1-(2-methylpropyl)-1H-imidazo[4,5-c]quinolin-4-amine,
 - 2-butyl-1-(2-methylpropyl)-1H-imidazo[4,5-c][1,8]naphthyridin-4-amine,
- 1-(2-methylpropyl)-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
 - 2-methylthiazolo[4,5-c]quinolin-4-amine,
 - 2-ethoxymethyl-1-phenylmethyl-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
 - 2-ethylthiazolo[4,5-c]quinolin-4-amine,
 - $\label{eq:def-amino-2-butyl-alpha-dimethyl-1H-imidazo} 4-amino-2-butyl-\alpha, \\ \alpha-dimethyl-1H-imidazo[4,5-c][1,5] \\ naphthyridine-1-ethanol,$
- 20 1-{2-[3-(3-pyridyl)propoxy]ethyl}-1H-imidazo[4,5-c]quinolin-4-amine,
 - 1-(2-phenoxyethyl)-1H-imidazo[4,5-c]quinolin-4-amine,
 - 1-[(R)-1-phenylethyl]-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
 - 1-{2-[3-(1,3-thiazol-2-yl)propoxy]ethyl}-1H-imidazo[4,5-c]quinolin-4-amine,
 - 1-[2-(pyridin-4-ylmethoxy)ethyl]-1H-imidazo[4,5-c]quinolin-4-amine,
- N-[3-(4-amino-2-methyl-1H-imidazo[4,5-c]quinolin-1-yl)propyl] cyclohexanecarboxamide,
 - 2-butyl-1-{2-[(1-methylethyl)sulfonyl]ethyl}-1H-imidazo[4,5-c]quinolin-4-amine,

N-{2-[4-amino-2-(ethoxymethyl)-1H-imidazo[4,5-c]quinolin-1-yl]ethyl} ethanesulfonamide,

- N-{2-[4-amino-2-(ethoxymethyl)-1H-imidazo[4,5-c]quinolin-1-yl]ethyl} propanamide,
- 1-[2-(methylsulfonyl)ethyl]-2-propyl-1H-imidazo[4,5-c]quinolin-4-amine,
 2-ethyl-1-{4-[(1-methylethyl)sulfonyl]butyl}-1H-imidazo[4,5-c]quinolin-4-amine,
 2-ethyl-1-[4-(ethylsulfonyl)butyl]-1H-imidazo[4,5-c]quinolin-4-amine,
 N-{3-[4-amino-2-(ethoxymethyl)-1H-imidazo[4,5-c]quinolin-1-yl]propyl}
 cyclopentanecarboxamide,
- 1-(2-methylpropyl)-6,7,8,9-tetrahydro-1H-imidazo[4,5-c]quinolin-4-amine,
 8,9,10,11-tetrahydropyrido[1',2':1,2]imidazo[4,5-c]quinolin-6-amine,
 4-amino-α,α,2-trimethyl-6,7,8,9-tetrahydro-1H-imidazo[4,5-c]quinoline-1-ethanol,
 2-hydroxymethyl-1-(2-methylpropyl)-6,7,8,9-tetrahydro1H-imidazo[4,5-c]quinolin-4-amine,
- 2-butyl-1-(2-phenoxyethyl)-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine, or a combination thereof.
 - 30. The method according to claim 22 wherein the dermal associated condition is selected from actinic keratosis, postsurgical scars, basal cell carcinoma, atopic dermatitis, and warts.

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31. The method according to claim 30 wherein the IRM is 2-methyl-1-(2-methylpropyl)-1H-imidazo[4,5-c] [1,5]naphthyridin-4-amine,
N-[4-(4-amino-2-butyl-1H-imidazo[4,5-c][1,5]naphthyridin-1-yl)butyl]-N'-cyclohexylurea,
1-(2-methylpropyl)-1H-imidazo[4,5-c]quinolin-4-amine,
2-butyl-1-(2-methylpropyl)-1H-imidazo[4,5-c][1,8]naphthyridin-4-amine,
1-(2-methylpropyl)-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
2-methylthiazolo[4,5-c]quinolin-4-amine,
2-ethoxymethyl-1-phenylmethyl-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
2-ethylthiazolo[4,5-c]quinolin-4-amine,

4-amino-2-butyl- α , α -dimethyl-1H-imidazo[4,5-c][1,5]naphthyridine-1-ethanol,

1-{2-[3-(3-pyridyl)propoxy]ethyl}-1H-imidazo[4,5-c]quinolin-4-amine,

1-(2-phenoxyethyl)-1H-imidazo[4,5-c]quinolin-4-amine,

1-[(R)-1-phenylethyl]-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,

1-{2-[3-(1,3-thiazo1-2-yl)propoxy]ethyl}-1H-imidazo[4,5-c]quinolin-4-amine, 1-[2-(pyridin-4-ylmethoxy)ethyl]-1H-imidazo[4,5-c]quinolin-4-amine,

N-[3-(4-amino-2-methyl-1H-imidazo[4,5-c]quinolin-1-yl)propyl] cyclohexanecarboxamide,

 $2-butyl-1-\{2-[(1-methylethyl)sulfonyl]ethyl\}-1\\H-imidazo[4,5-c]quinolin-4-amine,$

N-{2-[4-amino-2-(ethoxymethyl)-1H-imidazo[4,5-c]quinolin-1-yl]ethyl} 10 ethanesulfonamide,

 $N-\{2-[4-amino-2-(ethoxymethyl)-1H-imidazo[4,5-c]quinolin-1-yl]ethyl\}$ propanamide,

1-[2-(methylsulfonyl)ethyl]-2-propyl-1H-imidazo[4,5-c]quinolin-4-amine,

2-ethyl-1-{4-[(1-methylethyl)sulfonyl]butyl}-1H-imidazo[4,5-c]quinolin-4-amine, 15 2-ethyl-1-[4-(ethylsulfonyl)butyl]-1H-imidazo[4,5-c]quinolin-4-amine, N-{3-[4-amino-2-(ethoxymethyl)-1H-imidazo[4,5-c]quinolin-1-yl]propyl} cyclopentanecarboxamide,

1-(2-methylpropyl)-6,7,8,9-tetrahydro-1H-imidazo[4,5-c]quinolin-4-amine,

8,9,10,11-tetrahydropyrido[1',2':1,2]imidazo[4,5-c]quinolin-6-amine, 20 4-amino- α , α , 2-trimethyl-6, 7, 8, 9-tetrahydro-1H-imidazo [4,5-c] quinoline-1-ethanol, 2-hydroxymethyl-1-(2-methylpropyl)-6,7,8,9-tetrahydro-

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1H-imidazo[4,5-c]quinolin-4-amine, 2-butyl-1-(2-phenoxyethyl)-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine, or a combination thereof.

- The method according to claim 30 wherein the formulation further comprises a 32. preservative system and an emulsifier.
- The method according to claim 32 wherein the preservative system comprises 33. 30

methylparaben at 0.01 to 0.5% w/w of the formulation and propylparaben at 0.01 to 0.5% w/w of the formulation.

- 34. The method according to claim 32 wherein the preservative system comprises methylparaben at 0.01 to 0.5% w/w of the formulation and ethylparaben at 0.01 to 0.5% w/w of the formulation.
 - 35. The method according to claim 32 wherein the preservative system comprises iodopropynyl butylcarbamate.
 - 36. The method according to claim 32 wherein the preservative system comprises iodopropynyl butylcarbamate and one or more of methylparaben, ethylparaben, propylparaben, or phenoxyethanol.
- 15 37. The method according to claim 32 wherein the preservative system comprises iodopropynyl butylcarbamate, methylparaben, and ethylparaben.
 - 38. The method according to claim 32 wherein the preservative system comprises phenoxyethanol and one or both of methylparaben and ethylparaben.
 - 39. The method according to claim 32 wherein the preservative system comprises a preservative enhancing solubilizer.
- 40. The method according to claim 39 wherein the preservative enhancing solubilizer comprises diethylene glycol monoethyl ether, propylene glycol or a combination thereof.
 - 41. A method for delivering an immune response modifier (IRM) to a dermal surface, the method comprising the steps of:
- 30 selecting a formulation comprising:

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- (a) an immune response modifier selected from imidazoquinoline amines, imidazotetrahydroquinoline amines, imidazopyridine amines, 6,7-fused cycloalkylimidazopyridine amines, 1,2- bridged imidazoquinoline amines, thiazoloquinoline amines, oxazoloquinoline amines, thiazolopyridine amines, oxazolopyridine amines, imidazonaphthyridine amines, imidazotetrahydronaphthyridine amines, and thiazolonaphthyridine amines;
- (b) a fatty acid;

thiazolonaphthyridine amines;

- (c) a hydrophobic, aprotic component miscible with the fatty acid and comprising a hydrocarbyl group of 7 or more carbon atoms; and
- (d) a hydrophilic viscosity enhancing agent selected from cellulose ethers and carbomers; and

applying the selected topical formulation to the dermal surface.

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- 42. A pharmaceutical formulation comprising:
 an immune response modifier (IRM) compound selected from the group
 consisting of imidazonaphthyridine amines, tetrahydroimidazonaphthyridine amines, and
- a fatty acid; and
 a hydrophobic, aprotic component miscible with the fatty acid and comprising a hydrocarbyl
 group of 7 or more carbon atoms.
- 43. The formulation according to claim 42 wherein the formulation further comprises a preservative system.
 - 44. The formulation according to claim 42 wherein the hydrophobic, aprotic component has a hydrophilic lipophilic balance of less than 2.
- 30 45. The formulation according to claim 42 wherein the hydrophobic, aprotic

component has a pKa greater than 14.2.

46. The formulation according to claim 42 wherein the ratio of the hydrophobic, aprotic component to the fatty acid is 0.025:1 to 600:1.

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- 47. The formulation according to claim 42 wherein the combined weight percent of the hydrophobic, aprotic component and the fatty acid is 2 to 50.
- 48. The formulation according to claim 42 wherein the fatty acid is isostearic acid.

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- 49. The formulation according to claim 42 wherein the hydrophobic, aprotic component is selected from aprotic fatty acid esters, hydrocarbons of 8 or more carbon atoms, and waxes.
- 50. The formulation according to claim 49 wherein the aprotic fatty acid ester is isopropyl myristate, isopropyl palmitate, diisopropyl dimer dilinoleate, caprylic/capric triglyceride, cetyl esters wax, or combinations thereof.
 - 51. The formulation of claim 49 wherein the hydrocarbon of 8 or more carbon atoms is mineral oil or petrolatum.

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- 52. The formulation according to claim 43 wherein the preservative system comprises methylparaben at 0.01 to 0.5% w/w of the formulation and propylparaben at 0.01 to 0.5% w/w of the formulation.
- 53. The formulation according to claim 43 wherein the preservative system comprises methylparaben at 0.01 to 0.5% w/w of the formulation and ethylparaben at 0.01 to 0.5% w/w of the formulation.
 - 54. The formulation according to claim 43 wherein the preservative system comprises iodopropynyl butylcarbamate.

55. The formulation according to claim 43 wherein the preservative system comprises iodopropynyl butylcarbamate and one or more of methylparaben, ethylparaben, propylparaben, or phenoxyethanol.

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- 56. The formulation according to claim 43 wherein the preservative system comprises iodopropynyl butylcarbamate, methylparaben, and ethylparaben.
- 57. The formulation according to claim 43 wherein the preservative system comprises phenoxyethanol and one or both of methylparaben and ethylparaben.
 - 58. The formulation according to claim 43 wherein the preservative system comprises a preservative enhancing solubilizer.
- 15 59. The formulation according to claim 58 wherein the preservative enhancing solubilizer comprises diethylene glycol monoethyl ether, propylene glycol or a combination thereof.
 - 60. The formulation of claim 43 comprising:
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- (a) 0.001 to 5% w/w imidazonaphthyridine amine, imidazotetrahydronaphthyridine amine, thiazolonaphthyridine amine, or a combination thereof;
 - (b) 0.05 to 40% w/w isostearic acid;
 - (c) 1 to 30% w/w hydrophobic, aprotic component; and
 - (d) 0.01 to 30% w/w preservative system.

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- The formulation of claim 43 further comprising an emulsifier and a hydrophilic viscosity enhancing agent.

62. The formulation of claim 60 further comprising an emulsifier and a hydrophilic viscosity enhancing agent.

- 63. The formulation of claim 62 wherein the viscosity enhancing agent comprises a carbomer.
- 64. The formulation of claim 63 comprising:
 - (a) 0.03 to 3% w/w

2-methyl-1-(2-methylpropyl)-1H-imidazo[4,5-c] [1,5]naphthyridin-4-amine,

- N-[4-(4-amino-2-butyl-1H-imidazo[4,5-c][1,5]naphthyridin-1-yl)butyl]-N'-cyclohexylurea,
 - 2-butyl-1-(2-methylpropyl)-1H-imidazo[4,5-c][1,8]naphthyridin-4-amine,
 - 1-(2-methylpropyl)-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
 - 2-ethoxymethyl-1-phenylmethyl-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
 - 4-amino-2-butyl-α,α-dimethyl-1H-imidazo[4,5-c][1,5]naphthyridine-1-ethanol,
- 1-[(R)-1-phenylethyl]-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
 - 2-butyl-1-(2-phenoxyethyl)-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,

or a combination thereof;

- (b) 3 to 25% w/w isostearic acid;
- (c) 3 to 15% w/w hydrophobic, aprotic component;
- (d) 0.1 to 25% w/w preservative system;
- (e) 0.75 to 3.5% w/w emulsifier; and
- (f) 0.5 to 5% w/w carbomer.
- A method of treating a dermal associated condition, the method comprising a step of:

 applying to skin a formulation comprising an immune response modifier (IRM)

 chosen from imidazonaphthyridine amines, imidazotetrahydronaphthyridine amines,

 and thiazolonaphthyridine amines; a fatty acid; and a hydrophobic, aprotic component

 miscible with the fatty acid and comprising a hydrocarbyl group of 7 or more carbon

 atoms.

- 66. The method according to claim 65 wherein the ratio of the hydrophobic, aprotic component to the fatty acid is 0.025:1 to 600:1.
- 67. The method according to claim 65 wherein the combined weight percent of the hydrophobic, aprotic component and the fatty acid is 2 to 50.
 - 68. The method according to claim 65 wherein the hydrophobic, aprotic component is selected from the group consisting of aprotic fatty acid esters, hydrocarbons of 8 or more carbon atoms, and waxes.
 - 69. The method according to claim 68 wherein the aprotic fatty acid ester is isopropyl myristate, isopropyl palmitate, diisopropyl dimer dilinoleate, caprylic/capric triglyceride, cetyl esters wax, or combinations thereof.
- 70. The method according to claim 65 wherein the formulation further comprises:

a preservative system; and an emulsifier.

- 71. The method according to claim 65 wherein the IRM is 2-methyl-1-(2-methylpropyl)-1H-imidazo[4,5-c] [1,5]naphthyridin-4-amine, N-[4-(4-amino-2-butyl-1H-imidazo[4,5-c][1,5]naphthyridin-1-yl)butyl]-N'-cyclohexylurea,
 2-butyl-1-(2-methylpropyl)-1H-imidazo[4,5-c][1,8]naphthyridin-4-amine,
 1-(2-methylpropyl)-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
- 2-ethoxymethyl-1-phenylmethyl-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
 4-amino-2-butyl-α,α-dimethyl-1H-imidazo[4,5-c][1,5]naphthyridin-1-ethanol,
 1-[(R)-1-phenylethyl]-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
 2-butyl-1-(2-phenoxyethyl)-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine, or a combination thereof.

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72. The method according to claim 65 wherein the dermal associated condition is actinic keratosis, postsurgical scars, basal cell carcinoma, atopic dermatitis, and warts.

- 73. The method according to claim 72 wherein the IRM is 2-methyl-1-(2-methylpropyl)-1H-imidazo[4,5-c] [1,5]naphthyridin-4-amine, N-[4-(4-amino-2-butyl-1H-imidazo[4,5-c][1,5]naphthyridin-1-yl)butyl]-N'-cyclohexylurea,
 2-butyl-1-(2-methylpropyl)-1H-imidazo[4,5-c][1,8]naphthyridin-4-amine,
 1-(2-methylpropyl)-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
 2-ethoxymethyl-1-phenylmethyl-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
 4-amino-2-butyl-α,α-dimethyl-1H-imidazo[4,5-c][1,5]naphthyridin-1-ethanol,
 1-[(R)-1-phenylethyl]-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
- 4-amino-2-butyl-α,α-dimethyl-1H-imidazo[4,5-c][1,5]naphthyridine-1-ethanol,
 1-[(R)-1-phenylethyl]-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine,
 2-butyl-1-(2-phenoxyethyl)-1H-imidazo[4,5-c][1,5]naphthyridin-4-amine, or a combination thereof.
- 74. The method according to claim 72 wherein the formulation further comprises: a preservative system; and an emulsifier.
- 75. The method according to claim 74 wherein the preservative system comprises methylparaben at 0.01 to 0.5% w/w of the formulation and propylparaben at 0.01 to 0.5% w/w of the formulation.
 - 76. The method according to claim 74 wherein the preservative system comprises methylparaben at 0.01 to 0.5% w/w of the formulation and ethylparaben at 0.01 to 0.5% w/w of the formulation.
 - 77. The method according to claim 74 wherein the preservative system comprises iodopropynyl butylcarbamate.
- 78. The method according to claim 74 wherein the preservative system

comprises iodopropynyl butylcarbamate and one or more of methylparaben, ethylparaben, propylparaben, or phenoxyethanol.

- 79. The method according to claim 74 wherein the preservative system comprises iodopropynyl butylcarbamate, methylparaben, and ethylparaben.
- 80. The method according to claim 74 wherein the preservative system comprises phenoxyethanol and one or both of methylparaben and ethylparaben.
- 10 81. The method according to claim 74 wherein the preservative system comprises a preservative enhancing solubilizer.
 - 82. The method according to claim 81 wherein the preservative enhancing solubilizer comprises diethylene glycol monoethyl ether, propylene glycol or a combination thereof.
 - 83. A method for delivering an immune response modifier (IRM) to a dermal surface, the method comprising the steps of:

selecting a formulation comprising:

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- (a) an immune response modifier selected from imidazonaphthyridine amines, imidazotetrahydronaphthyridine amines, and thiazolonaphthyridine amines;
- (b) at fatty acid;

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(c) a hydrophobic, aprotic component miscible with
the fatty acid and comprising a hydrocarbyl group of 7 or more carbon
atoms; and

applying the selected formulation to the dermal surface.

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C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category •	Citation of document, with indication, where appropriate, of the rel	levant passages	Relevant to claim No.
X ·	CHOLLET J L ET AL: "DEVELOPMENT TOPICALLY ACTIVE IMIQUIMOD FORMUL PHARMACEUTICAL DEVELOPMENT AND TO NEW YORK, NY, US, vol. 4, no. 1, January 1999 (1999 pages 35-43, XP000900717 ISSN: 1083-7450 abstract page 36, column 1, paragraphs 3, page 40, column 2 -page 41, columparagraph 2 page 42, column 2, paragraph 4 -page 42, column 2, paragraph 4 -page 42, column 2, paragraph 4 -paragraph 2	LATION" ECHNOLOGY, 9-01), 4 nn 2,	1-3,7,8, 10-12, 18,22, 25, 27-29,41
X Furti	ner documents are listed in the continuation of box C.	X Palent family members	are listed in annex.
"A" docume consid "E" earlier of "Illing d "L" docume which citation "O" docume other r "P" docume later th	nt which may throw doubts on priority claim(s) or is câed to establish the publication date of another n or other special reason (as specified) ant referring to an oral disclosure, use, exhibition or	cited to understand the prin- invention "X" document of particular releva- cannot be considered novel involve an inventive step with "Y" document of particular releva- cannot be considered to invi- document is combined with	onflict with the application but ciple or theory underlying the unce; the claimed invention or cannot be considered to the the document is taken alone unce; the claimed invention olve an inventive step when the one or more other such docuping obvious to a person skilled one patent family
2	2 April 2003	07/05/2003	
Name and n	nailing address of the ISA European Patent Cifice, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tet. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Authorized officer Houyvet, C	

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ation) DOCUMENTS CONSIDERED TO BE RELEVANT	
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WO 00 40228 A (3M INNOVATIVE PROPERTIES CO) 13 July 2000 (2000-07-13) page 1, line 5-15 page 1, line 23 -page 2, line 7 page 4, line 10-15 page 5, line 5 -page 27, line 15 page 28 line 9 -page 31 line 2	1-83
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example 10 US 6 110 929 A (MARSZALEK GREGORY J ET AL) 29 August 2000 (2000-08-29) column 1, paragraph 2 column 6, paragraph 5 -column 7, paragraph 1	1-83
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US 4 595 586 A (FLOM MERLYN G) 17 June 1986 (1986-06-17) column 2, paragraph 2; example 2	1-83
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Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This Inte	emational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. χ	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
	Although claims 22-41 and 65-83 are directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged effects of the compound/composition.
2.	Claims Nos.: because they relate to parts of the international Application that do not comply with the prescribed requirements to such an extent that no meaningful international Search can be carried out, specifically:
з. П	Claims Nos.:
٠. ا	because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)
This inte	ernational SearchIng Authority found multiple inventions in this international application, as follows:
1.	As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3.	As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4.	No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the daims; it is covered by claims Nos.:
Remark	on Protest
	No protest accompanied the payment of additional search fees.

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